STRUCTURAL STUDIES OF SOME NATURAL POLYSACCHARIDES

A THESIS

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Cortified that the thouse antitled, "STRUCTURAL STRUCTURAL STRUCTU

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Security Section Services

The dispostation entitled, "STRESTURAL STURES of SCHE MATURAL POLYSACCHARDES", deals with the isolation and chemical examination of polygoochasides from the seeds of <u>Signature injula</u> and <u>Phaspolus surge</u>, some constituents from the seeds of <u>Dausus</u> Sample, and free unter colubic managedarides from the flowers of <u>Hama, unilasinalizas</u>. The thouse has been divided in to five chapters.

The Chapter I is of introductory nature and describes the wide importance of natural products and a brief secount of different cleases of compounds, i.e. polysapphurides, starols, flavornoides and free sugars.

The Chapter 11 deals with the isolation and structural elucidation of neutral water soluble polysoccharide from the seeds of Zizyphus jujubs.

The Chapter III describes the isolation and structural elucidation of a water coluble neutral polysecoharide from the seeds of Phaseolus sungs.

The Chapter IV. is divided into three Sections (A), (3) and (C), deals with the implation and elucidation of chamical structures of a sterol and two flavonaids from the seeds of Dammus.

The Chapter V and the last chapter forms the subject metter of chemical examination of free water soluble menosaccharides from the flowers of Linux unitationisms.

A imief review of uptodate literature on chandral exemination of selected plants, has been described respectively in each concerned chapter.

The work represented in the thesis has been carried out in the Chanical Laboratories of Dayanand Vedic Post-graduate Callege, ORAL, under the Supervision of Dr. C. S. Hiranjan, D.Phil., F.L.C.S., Department of Chamistry, Dayanand Vedic (P.C.) Callege, Chi.

A Wild's demonstry of the emplice work has been submitted percotrately along with the thrests, scoopeding to the peoplements of Cartinopeds for No.D. degree of Europhidani University.

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GHBARCAL LABORATORIES, D.V. (P.G.) COLLEGE, ORAZ-282001. Kekama Rani Gupta (KSHAMA RANI GUPTA)

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GRAVIER - A

DATA MODELLICA

Prom the time immemorial dispuses have been the cause of mark before human sufferings. Plants have been proved beneficial from the earliest times in curing various eliments and dispuse.

A continued search for modicinal plants during the last several conturies has given us a built of modicinal plants which are of great use in the treatment of dispuses and promotion of health.

definite chemical constituents in them. Medicinal plants may sometimes contain some tomic substances. That is why, the use of plants in their natural states is not proper and isolation of active principles in pure state from medicinal plants is very essential. The chemical investigation in the field of Netural Froducts gained pass with the advent of the use of developed new techniques like chromatography, spectrometry 2,2,4 and verious physico-chemical methods. These modern techniques are useful to isolate; active organic compounds even when they are present in a very small quantity.

There are many families and genero in the vest flore world which have high medicinal values and have not been investigated as yet for their active principles. The new researches on these plants may prove quite beneficial to cure human eliments.

Various chamical constituents obtained from the plants are classified into mapy groups. A brief account of the review on the

classes of compounds investigated from the plants, which have been incorporated in the present thouse is given below

1.2 * Polysocchemides.

2.3 - Stanola.

1.4 - Playonoide.

1.5 * Page gugage.

1.9 POLYSACISMUSS

Polysocchardes are most important component of all living organisms and highly distributed among the higher order of land plants and see woods. They are present in Sungi empireleton of insect and crustopous, in the copules of microorganism, in certilogs, in emimal joint fluids etc.

Polysuccharides are magranolocular compounds, composed of several monosocharide units, usually linked through exygen to give complex composition. They are hydrophillic colloids of high mole-cular weight, some completely soluble in water, other swall and absorb considerable amount of water without dissolving.

Gues and maniloges are complicated polysacchoride polymers and differ in the respect that the former are characterised as plants emudates while the latter are isolated from various plant organs by extraction with water.

Plant gums and mucilogos have been known and in use since very early times, reference being made to them in the sible; and they seem to have been of communcial value for several thousand years, especially in India, Asia, Agrica, Austrolia, and Ching.

There is no agreement as to the exigin of gum emudates, but they play an important role in the physiology of plants, in animal and microorganism as surface material and regarded as food reservore. In such the same manner as starch in many plants and glycogen in animals or as agent for holding water. The plant is ballowed to synthesize the gum emulates in arrier to seal off the infracted section of the plant and prevent further invesion of the tiesue. The plant and prevent further invesion of the tiesue. The gume may be, it is responsible to ballowe that gum anniates are formed by some type of enzymic polymerisation and not by disset polymerisation.

Ours and mucileges are used in while range of industries like commetties $^{10_{a},1_{a},12}$, phormacy $^{13_{a},14}$, textiles $^{10_{a},1}$, adhesives 17 , food products $^{10_{a},19_{a},20}$, paper $^{21_{a},22}$ and in many other fields.

A polysecthoride is isolated from the plant by extraction with cold or hot water, water containing a little acatic acid and the precipitation of the soluble portion with the empess of ethanel. The polysecthoride is purified to remove the inorquaic ions and proteinous impurities. By reported precipitation with ethanel from acidified equapus solution.

The homogenoity of the polymercharide is checked by drace tional precipitation, some electrophoresis 25,25, and sectylation and descriptories. A mixture of polymercharides can be espansive ever a colluious tolumn 25,25, while the soldie polymercharides may be frestionated as their complement. Jos ambience column 25, 25

also used effectively for the fractionation but the methylated gums are separated over alumino³². Electrophoretic separation of polysectoration has been achieved mainly in herate buffer^{33,34}, but seetate buffer^{35,36} and citrate buffer³⁶ have also been used. With the help of membranes or filters of desired percenty³⁷ polysecharides may be fractionated. The unumbed polyseccharides of the mixture may be destroyed with specific engages³⁰ followed by denoturation of ensure with heat, alkali and alcohol. The fractionation of polyseccharides may also be achieved by gel filtration³⁰ and molecular slave⁴⁰.

The purified polysaccharded is subjected to preliminary determination of lights, san content, methonyl, acetyl, primary bydroxyl and carbonyl groups and they are estimated after the determinate of nitrogen, sulphur, phosphorus and halogens which may be present in the polysaccharde.

The optical rotation of the polysaccharide is measured by means of usual polarimeter ⁴¹ or photosicetric spectropolarimeters ⁴² The configuration of phycosidic linkage in oligosaccharides can be correlated to optical rotatory power by applying Hudson's rule of isomotation ⁴³.

The molecular weight of the polymarcharide having terminal reducing group can be determined by estimating it with c¹⁴ labelled sodium cyanide⁴⁴, sodium hypotedite⁴⁵, ferricyanide⁴⁶, and periodete exidation studies⁴⁷. Physical methods like viscosity⁴⁸, light scattering⁴⁸, cometic pressure⁵⁰ are also used to determined the molecular weight of the polymarcharide.

The hydrolymis of the polymorphoride with mineral saids under different conditions provides information regarding the nature of linkages present between sugar modeties. The complete edd hydrolysis of the polysecheride results in the liberation of monosaccharides which can be esperated by paper 22 or column chromotographic 52 techniques. They are identified by their Re values. co-chromatography with authentic camples, melting points and by proporing their drystelline derivotives. Partial acidic hydrolysis with dilute mineral ecids (0.0: * 0.11) results in decredation of the polyseconomide into less complicated melecules which can comily be identified. Oligosaccherides, obtained by partial hydrolysis, can be separated by paper chromatography and their etructure is determined by the usual process of methylation, followed by the bydrolysis and identification of methylated sucare, periodate oxidation and enzymic bydrolymia. Enzymic degradation 33 provides various information about the polyeocharide.

The sugars may be quantitatively estimated by microvelumatric method, spectrophotometric method or colorimetric method. Accountly as extensive use of gas liquid partition chromatography 54-56 in the separation and estimation of sugars has been reported.

The polysoccharade is subjected to paradiate emidetics to obtain the information regarding the nature of end groups and types of glycosidic limitage present. It has been observed that the 1,2-dial groups in $1 \Rightarrow 2$ or $1 \Rightarrow 4$ limited and 1,2,3-trial groups in the $1 \Rightarrow 6$ limited subjectively, liberating one make of female

acid but the units howing 1 >> 3 linkages with no l_s2-diel system are not effected. Thus by determining the consumption of periodete and amount of femic ecid liberated, various informations regarding the etructure may be obtained.

The methylation studies serve the valuable information regarding the types of linkages between sugar moleties in a polyseccharide. The method consists in the methylation of the polyseccharide followed by hydrolysis to give methylated sugars. The nature and the quantitative determination of the methylated sugars provide information on the relative proportions of non-reducing and groups, the degree of branching, the type of interchain linkages and the nature of the main chain linkages in the polyeocheride. Nothylation is usually carried out by means of Howorth's method 55 followed by Purdle's method ". The methyleted product is bydeelysed in two steps, first the methonolic hydrogen chloride or with 80-90% formic ocides and finally with the mineral acids. The methylated sugars are separated on paper and identified by their R_{TMG}^{-62} values, optical rotations and melting points of their crystalline derivetives. The methyleted sugars are quantitatively estimated by titrating them with alkaline hyperodite or by coloria metric method. Those polysaccharides which are soluble in disethyl sulphoside, may be very efficiently methyloted in fewer steps by using so thyl iodide and cilver omide.

In the present theels, the chemical emminetion of the complex water soluble polyeaccharides, a galactomerman limited from the seeds of Zimphra jujube and a galactomerman limited from the seeds of Phaneolius annot have been described in Them. It nespectively.

They are crystalline compounds, and contain an alcoholist group. The structure of the stances are based on the 1:2 eyelo-pentenophananthrone skeleton. The stances give characteristic Liebonnann-Burghard reaction.

The plants have veriety of closely related starols called phytostorols. They occur in the plants in free state or as estars of higher fatty scids or emotions as glycosides called staroline. Heny of them are isolated from the unexpendicable portion of alls and fats. The well known phytostorols are stigmateral. β -sitestorol and expectance.

The sterols are found to be physiologically important substances, play various roles in life process and have great importance in animal metabolism, hermones, co-enzymes, bile acids and provitamin-D.

The investigator has been able to isolate a \$-aitestard from defetted matters of the soods of Deucus carets. The chamical study of this sterol has been described in Chapter IV of the thesis.

1.4 PLANOSTOTOS

Flavonoids form the Largest group of naturally ecomoring outside to page the group of naturally ecomoring outside the compounds as page and a large scale of $C_3 = C_3$ canbon shale to a solic two because where joined by a taxon cambon that which in femore into a various slaves of flavoures compounds a figure a

isoflavonos, flavonols, flavanononols, dihydroxyflavonoles, flavanonos, isoflavononos, chalkonos, dihydroxychalkonos, auronos, anthocyanidins and laucoanthocyanidins, differ from one another only by the state of emidation of this 3-C-link.

Flavonoids are propert in plants in the free state at unil as in the ferm of glycosides, containing either sugars on more than one hydroxyl grouper disaccharide (bioside) and trisaccharides.
Nest plants contain more than one glycoside of any aglycose.

It is supposed that flavores protect plants from hazminical ultraviolet radiations or from less of important metastals by autoomidation and one is tempted to believe physiological functions of the flavored pignents besed upon their colours are related. to the role of flavors in reproduction ⁶⁴. These compounds were found to be of great medicinal importance as becteriostatic ⁶⁵ and importance as becteriostatic.

1.4.1 PLAYONES AND PLAYORING

The flowenes and floweness are naturally colouring matters. Their structure is besed on that of 3-phonyl-4-chromene. The flowenes and floweness differ in the respect that latter has a hydroxy group at position ~3. The besic skeleton of flowene and flowenes hay be respected as.

Playmol skalaton

Flavone, akalatan

The structure of those compounds was not proporty clusted dated until 1891 elthough Morin was isolated as early as 1814. In 1891 Hertaig to posted the structure of quameria. Afterwards (1893) the structure of chaysin was determined. Today nearly one bundred flavones and flavoness have been isolated, the letter class making nearly two-third of the total 60.

These compounds occur naturally in free state or as glycosides. The position occupied by a sugar unit in glycoside linkage, plays an important part due to which a glycoside embibits difference in proporties as solubility and capacity to form complement with metals. Unlike anthogyanine in which the sugar residue is usually present at position 3 and 5, the sugar molecy in flavones and flavones is generally attached to a hydroxyl group at position 3 or 7.

These compounds have been found to be highly physiologically estive. The flavonol physoside mutin has been described for its therepoutic properties. The importicidal action of polyhydromy flavones and their ethers and the estion of flavones on included enzyme eyeten⁶⁹ heve been etudied.

The author has been able to isolate a flavone glycoside and a flavonel compound from the seeds of <u>housels supple</u>. The chamical study of those colouring substances has been described in Chapter IV of the themis.

1.5 FRUE HERER STREET, STREET, (CARSTRYTEATER)

Carbohydrates are an important class of naturally occurring substances and are found universally distributed same plants, animals, and micro-erganisms. The name carbohydrate arose from the fact that the first compounds of this group to be studied ware found to have an empirical formula $C_{\chi}(H_{\chi}O)_{\chi}$ and were believed to be hydrates of carbon. Since that time, however carbohydrates which do not have hydrogen and oxygen present in the proportion to form water $(a, g, xhannose, C_{\chi}H_{\chi}O_{\chi})$ has been discovered, and other carbohydrates containing nitrogen and sulphur are also known. Although it is difficult to define such a heterogeneous group, the carbohydrates may be thought of as polyhydroxy aldehydes or ketones and derivetives of them.

Catabolism of carbohydrates provides the major share of the energy requirement for maintenance of life and performence of work. The metabolism of carbohydrates is of central importance to organism, individually and collectively. Besically all organic food-stuff are ultimately derived from the synthesis of carbohydrates through photograthesis. Carbohydrotes are divided into three basic categories :
Nonesaccherides, oligosaccherides and polysaccherides. Nonesaccherides have three to nine, usually either five or six, carbon stans and contain only one aldehydic or ketonic functional group. The oligosaccherides are oligomers of anesaccherides linked by formstance of glycosidic linkages. These generally contain two to eight or ten menomeric units. Folysaccherides are frequently molecules of great size and may have molecular weights of many million. They contain more than ten menomeric units.

In the present thesis in Chapter I the study of water soluble monosaccharides from the flowers of Linux unitationisms is incorporated.

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CHAPTER - 18

A REST WATER SOLUBLE POSTERCORARDE

AND HE SEED OF

Liels. The subject matter of this Chapter is isolation and characterisation of a neutral nature soluble polymercharide from the seeds of Zizyphus jujuba.

The plant limphus juiche Lank. commonly known as Jerberi (Indien jujube). The cultivated tree is called "Pawandi" or "Sundiber". This plant belongs to the family Shannesceet, a shrub or nederate-size tree, almost evergreen, usually armed. Young branches and flowers densely tementose. Leaves variable, i — inches long, evate-oblong or sub-orbicular, obtuse or scute, course or serulate, dark green and glabrous above, clothed beneath with dense pale-coloured tomentum. Frickles solitary and straight or in pairs with one of then shorter and recurved, rarely usinting-flowers greenish-yellow scmowhat jostid, arranged in short axiliary, subsecuile cymes. Calyx glabrous within. Futals clawed, with an oblong hooded lamins. Jisk 10-lobed. Ovary 2 called, styles 2, connets to the middle. Drupe & & inches or longer, globese oblong or evoide, orange or red when ripe. Stone 2 - called, with a hard thick bony shell.

Indigenous and neturalized throughout India and in Coylon, which and cultivoted, also in Tropical Africa, the Malay Archipelage, Chine and Australia.

Various parts of the tree are used medicinally². The fruit is largely esten by notives and it is such valued in times of earcity and it was consider/to purify the blood and aid digestion. Not describe used in fever and as a pender applied to old wounds and ulcome. But considered to be a mandy in discussor. The

who of Egyption wood in ancient civilization is reviewed, and the chilities of many these species to have registed termite attack was studied. This specimen is 3000 * 4000 years old. The oldest Egyption wood belongs to the genus Lizyphus³.

11.2. The Described of Charles Examination of this plant in the Literature is Described of given below :

Convo		Plant species	Conetltuents		
1.0	21 s yphu s		Vitania G Contont		(1936)4
2.	21270000	•	Anthroquinone derivetive		(1989) ³
3.	Jujuba (Florida gross)		Garotane and aterrisic edid contant		(1949)9
4.	Jujube (Kamaa)		lysine. espartic ocid glysine.espargi glutamic ecid & galectose	ne ,	(1969) ⁶
•	Zizyphus	Taleni (Balkat/ (Balco)	iignin end	Mod (Falli- poine)	(1932)
Ú,	Zizyphuo				(1939)0
70	21syphus		Starch mois- ture (8.4%, esh contont 1.63%, starch proper 1.22	Socio	(3949)9
9.	Ziz yolmo	Joomerio	Detargent		(1951)10
9.	ZLeyphus	J_omorio	Jungie acid		(2000)
0.	Zizy hus	Nylophore .	Sutudinic ocid		(1901)**
L .	Zisychue	Mylophyrus	Tonnine & oleenolic agid	Prules	(1969)23
20	<u>Aspino</u>	hylophyrus	(-)-Lougoantho- eyanin	Part No.	(1920)24
		Concollo	Two new boole poptide liny- phine; linysti- nine (betwinte		(1969) ₇₉

Genus	Plant species	Constituento Parto	
		ecid, 0-glucoge,	
		> fructose.	
LA. Lington	Cenoplia	Constitution of Root & Zizyphinine Bark	
19. Zigyphus	Kauritiene	Two poptide alkaleids Mauritine (A) & Mauritine (A)	(1972)17
16. Lizyphus		Ester solu- ble carbo- hydrates.36.1% fructose. 22.3% D-plucose.14.8% oligosescheride. 1.4% arabicose. 2.5% gologurones.	(1960) ²³
17. Zieyphas	Acondorie	No. K. Ca. No. Po. Al. Cu. end En trace mineral constituents	(1970) ¹⁹
16. Zizyphue	Vulgaria	Forty unid and Jark routh noids from other outrest	(1934) ²⁰
19. Zlayphus	Vulgario	Anaosthotics Leaves	(1841)51
20 • 21 gyphus	Vulgazio	Chinese des Sanda (extracted of β) and factor β and β a	(1936)
ll. Zlayphue	Wignels	Setulinic acid Seeds (C ₃₀ H ₄₆ O ₂)	(1946) ²³
2. Azyjtas	Jujubo	Loucocyanidin Hark & Loucopelaryenidin wood and Setulinic & Cossothic acids	(1961) ¹²
D. Zasyches	Jujubo	Caryl elcohol, Leaves Alkaloids, Protopine & Sasbaria	(1930)
4. 227722	Adus	Ortin Laws	(1968)25

Gold Michigan	Control of the sense of				
	140	Plant Species	Gonotituents	Astr.	
23	Zisyphus		five alkaloids of 13 membered cyclopoptide alkaloidal ring structure	20000	(1970)26
	24 syphus		Tenning, Anthre-	Pruits & Looves	(1960)27
	44.5 young		Carbohydrates Carotane Gaonins Elevone glyce Gides, seponins Lipids, resins and surilage		(1969)20
20.	21syphue		Oil, contained olais, lineleis, arachidis and behanic acids.	loods	(1953)29
29.	ZIBY AND	Jujoba	Secontial amino :	a0.0s	(1969)30
30.	Z18yphuo	Jujubo	Sapegemin (Spalin lectors)		(4970)31
	Lizyphus	Jujuba	Sapomin(Jujubosis D _i structure elus dation by cambon- nuclear megnetic resonance		(1978) ³²
	Z2zyphus	Jujuba	Attach		1969/32

A sumber of chemical compounds have been already reported in the above literature, but no attempt has been made for the isolation and structure elucidation of polymercharicae of <u>Limbus luids</u>. Second of the medicinal and industrial values of the plant, it was considered worthwhile to isolate and establish

the structure of their polysoccharide leolated from the seeds of \underline{z}_* jujubs.

MASS STRUCTURAL BLUCTDATION OF BRUTRAL BATER SOLUBLE FORMS

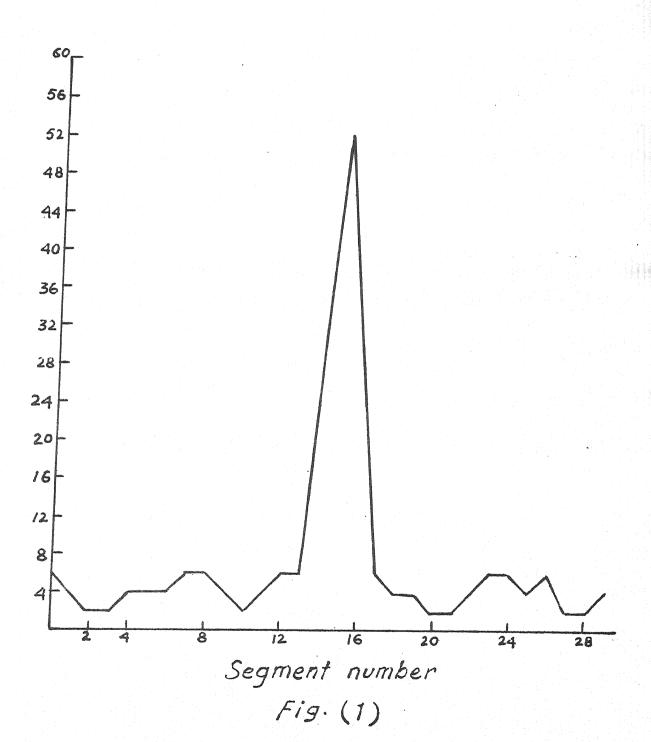
Bears Wa occise of

The polymorcharide was isolated from the defetted mede of Z. jujuba, entrocting with unter (1% aperic acid) and procipitating with ethanol. The polymorcharide was purified by repeated precipitation with ethanol to get a white dibrous sucilage with minimum ash content (0.6%). The hemogeneity of the polymorcharide was checked by.

- (1) Practional procipitation.
- (11) Zono electrophoreois.
- (111) /cotyletion and descetylation.

The polyeaccharide was dissolved in veter and separated into two fractions by fractional procipitation with different volumes of ethanol. Note the samples were analysed quantitatively by the method of Hirst and Jones 45. The results were essentially identical showing the homogeneity of the polyeaccharide.

The polymorcharide was acctylated with acetic anhydride by the usual method to give the acetylated product, $[\kappa]_0^{21} = 90^\circ$ (in acetone, C, 1.1%). Seacetylation of the product gave a polymorcharide having the same optical activity/as the original one. This confirmed the homogeneity of the polymorcharide.



Another portion of polysocharide was separated by some electrophoresis in borsto buffer (pi 9.3). The paper chromatogram was cut into 1.0 em segments, which were numbered consecutively from anodic end down to cathodic end. Each segment was eluted with distilled water, treated with phonol-sulphuric soid reagent and the absorbance of characteristics orange yellow colour was measured in a Klett-Summerson photoelectric colorimeter, using filter No. 30. A plot of the absorbance against segment number showed only a single sharp peak (Fig. - 1) indicating the polysaccharide to be homogeneous.

The polysaccharide was slowly soluble in water, [4] = 91.2° (in water, G. 0.6 g per 100 ml of solution), ash content 0.5%. The polysaccharide was found to be free of nitrogen, sulphur, and halogens. The methoxyl, uronide and acetyl percentage were found to be negligible.

The complete acid hydrolysis of the polysoccharide with authoric scid followed by the paper chrometographic analysis of the hydrolysate revealed the presence of two sugars. --galactose and --mylose. The identity of the sugars was confirmed by their specific optical retations, proparation of their crystalline derivatives and co-chrometography with authoric samples.

The quantitative estimation of monosecthoride components by periodate exidetion, taking ribose as a reference sugar, showed that galactose and mylose are present in the moler ratio, is in the polyeacheride. The graded bydrolymic of the polyeacheride with 0.05% sulphuric acid and subsequent paper chromatographic

analysis of the hydrolysotes, taken out at various intervals, revealed that Degalectose was liberated first followed by the liberation of Denylose. This shows that meet of the sylose units are linked together forming the beskbone (main chain) of the ployescharide and galectose units are linked as terminal groups.

The polysaccharide was methylated first by Hawarth method using dimethyl sulphate and alkali⁵⁷ followed by Fundic's method⁵⁸ with methyl iddide and silver exide to give/ a methylated product, [2] = 36° (in chloroform), G, 1 g, per 100 ml of solution),

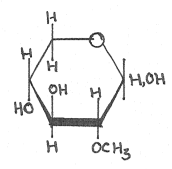
Ome, 44.6%. The complete hydrolysis of the methylated polysacchemide and paper chromatographic analysis of the hydrolysate in

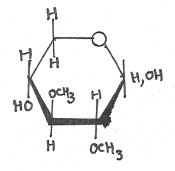
Solvent A, revealed the presence of four methylated sugars. The
methylated sugars were separated on a preparative scale by chromatography on Whatman No.3 filter paper. The following methylated
sugars were identified.

- (I) 2-0-omthyl-0-mylone.
- (II) 2,3-01-0-methyl-0-mylone.
- (III) 2,3,4-Tri-O-mothyl-D-mylose.
 - (IV) 2,3,4,6-Tetro-O-mothyl-3-galactoce.

Hethylated sugar, I, had R_{IRG} value in solvent A, 0.39, $[K]_{0}^{25} = 24^{\circ}$ (in water, C, 25), n.p. 130-32°. It formed 2.0-methyl-D-mylose smillde, n.p. 123-24°, $[K]_{0}^{25} + 213^{\circ}$ (in ethyl acetate, C, 0.65). Its discretate, 2-Omethyl, 3.4-discretate had n.p. 73-77°, $[K]_{0}^{25} = 38.5^{\circ}$ (in chloroform, C, 2.55). Thus the shows observations confirmed that the methylated sugars, I, is 2-0-methyl-D-mylose.

Nethylated sugar, Il , was obtained as a syrup, $A_{\rm DEC}$ in solvent A, 0.76, $[<]_0^{20}$ + 22.2 (in water, C, 4.30)), Ole, 34.5).





11

III

IV

It formed anilide derivative, 3,3-di-O-methyl-D-mylopyranesyl anilide, m.p. 130° , $[<]_{0}^{21}$ + 192.3° (in ethyl ecetate, C, 0.223%), which shows that the methylated sugar, II, is 2,3,-di-O-methyl-D-mylose.

systellised, $R_{T/SG}$ in solvent A, 0.92, $[K]_{S}^{16} + 19.2^{\circ}$ (in water, C, 0.39). On treatment with ethenolic enline it gave 2.3.4. tri-0-methyl-0-mylopyranesyl enlide, m.p. 94-96°, $[K]_{D}^{22} - 83^{\circ}$ (in ethenol, C, 26). The sugar in this fraction was thus identified as 2.3.4-tri-0-methyl-0-mylose.

Hethylated sugar, IV, $n_{\rm IMG}$ in covert A, 0.90, $|\mathcal{C}|_0^{25}+12e^6$ (in water, C, 0.6%), n.p. 72-73°. On trestment with ethenolic entline gave . 2.34.6-tetre=0-methyl-1-phenyl-3-galactosylamins, n.p. $180-90^{\circ}$, $|\mathcal{C}|_0^{25}=80^{\circ}$ (in acotene, C, 1%). Therefore, the identity of methyleted sugar, IV, is established as 2.3.4.6-tetre=0-methyl-3-galactose.

The quantitative estimation of methylated sugars, by the method of Hirst and Jones 60 showed that the sugars, I. II. III. and IV were present in the melecular ratio 5:0:213.

The appearance of 2,3,4,6-tetra-0-methyl-3-galactose, IV an 2,3,4-tri-0-methyl 3-mylose, III, on hydrolysis of methylated polysoccharide indicateSthat all galactose units and mylose (2 units) in the polysoccharide occupy terminal position as non-reducing and groups. A large proportion of, II, 2,3-di-0-methyl-1-mylose (8 moles) idmicates that the backbone of the polysoccharide consists of D-mylose units limited through 1 >> 4 linkages.

Detection of 2-0-mothyl-D-myloso, I, (5 males) shows that five mylose units in the main chain per repeating unit of the polysecheride are linked at position 3 in addition to 1- and 4-positions.

Determination of terminal groups by periodate exidation and subsequent titration of formic acid liberated corresponds to 0.2066 moles of formic acid per 100 g of the polysaccharide. On the basis of methylation studies, the simplest repeating unit of polysaccharide, is supposed to consist of 18 sugar moleties of which 3 units of galactose and 2 units of sylose form terminal groups, considering such a repeating unit, the terminal groups were found 28.33% as determined by periodate exidation studies, which is in close agreement to that revealed by mothylation studies (27.71%).

During the periodate exidation studies, the exidiced polysected was taken out from the reaction mixture after 72 hours and hydrolysed after destroying the periodate. The paper chromestographic examination of the hydrolysete showed that the presence of sylose was quite prominent, while no galactose could be detected. The paper chromatography of the hydrolysete of the exidiced polysectharide taken out from the reaction mixture after 96 hours showed the absence of both the sugars. It reveals that galactose units were completely exidiced within 72 hours, whereas sylose units were exidiced only after 96 hours. The considerable difference in the rates of exidation of the component appears is due to storic effect resulting from the branched structure of the polysectharide. The present knowledge, however, indicates that this phenomenes is mist likely due to cyclic sected formation.

The pertial acid hydrolysis of the polysaccheride followed by paper chromotographic separation on preparative scale affected six oligosaccharides which were detected as follows :

- 1. 3²-β-Mylobiocyl mylobioco (1 → 4-0--β-W-mylopyrmocyl
 -1 → 3 -0-β-W-mylopyrmocyl-1 → 4-W-mylopyrmocyl or
 3²-β-Mylocyl mylotrioco (0-β-Wmylopyrmocyl-(1 → 3)

 0-β-W-mylopyrmocyl (1 → 4)-0-β-W-mylopyrmocyl(1 → 4)-W-mylopyrmoco).
- 2. 3²-β-Nylosylxylobioso (-0-β-N-xylopyransyl-(1 → 3)-O-β-N-xylopyranosyl--(1 → 4)-D-xylopyranoso).
- 3. Nylotzioso(-0- β -0-nylopyzonosyl-(1 > 4)-0- β -0-nylopyzonosyl-(1 > 4)-0- β -0-nylopyzonoso).
- 4. Rhodymonobiose(«C» β «C»xylopyranosyl»(1 ⇒ 3)»C» «C»xylopyranose).

Gligoseccharide (1), $|\zeta||_{0}^{21} = 57.6^{\circ}$ (in water, C, 5%) found chromotographically sure in two solvents F and S. The complete edd hydrolysis followed by paper chromotographic analysis revealed the presence of only sylose units in the eligoseccharide. The molecular weight 550.3, of the eligoseccharide of corresponded to a tetraspectacide of pentoses. Partial acid hydrolysis of tetraspectacide gave eligoseccharides, $3^{2} * \beta * mylosyl * mylobioss*.$

mylotriose (-0-3-0-mylopyranosyl-(1-)4)-0-3-0-mylopyranosyl- $(1 \Rightarrow 4)=0-\beta =0$ -sylopyzanose), zhodymenablese and sylobiose, carrespended, the the oligopaccharides (2), (3), (4) & (5) respectively. A (1 -> 3) linkage in the oligopaccharide was also confirmed by periodate oxidation. The consumption of 5.2 moles of metaperiodate. with the liberation of 2.16 moles of formic acid per male of the oligosecharide. Had, all the sugar modeties in the tetragache* ride been linked by (1 > 4) linkeges the tetrasecheride would have consumed 6 moles of periodate instead of 5.2 moles. The hydrolysis with the enzyme emulsin and the negetive rotation indicated that the sylose units in the eligosaccharide were linked through β . linkages. On the basis of these experimental evidences, the oligosaccharides have been identified as (1 → 4)-0-3 -0-gylppyranosyl- $(1 \rightarrow 3)$ -0- β -0-xylopyranosyl- $(1 \rightarrow 4)$ -0-xylopyranose, 1.e. $3^2 - \beta$ -mylobiosylmylobiose or $0 - \beta$ -0-mylopymonosyl-(1 ->3)-D-mylopyranosyl- (1 -> 4)-O- \$-D-mylopyranosyl-(1 -> 4)-O-mylopyranosa. i.e. 33- A-mylosylmylotziose. Fig. - 2(a) and 2(b).

Oligosecharide (2), m.p. 222° , [\checkmark] $^{\circ}_{D}$ - 51° (in water, C, 2.9%), was chromatographically pure in solvents F and 3. The molecular weight 420 corresponded to a trisocharide of pentoses. Acid hydrolysis of the oligosecharide yielded only mylose. The anomeric configuration of non-reducing mylose units were found to be $^{\circ}\beta$ by enzymic hydrolysis and negative rotation. Partial acid hydrolysis yielded, mylobiose, rhodymensbicse, corresponding to eligosecharides (5) and (4) respectively and mylose which were identified by co-chromatography with the authentic samples. Pariodete oxidation studies revealed the consumption of 4.3 moles of metroperiodete with the liberation of 2.1 moles of formic poid.

Fig. - 2(a).

Fig. - 2(b).

Fig. - 3.

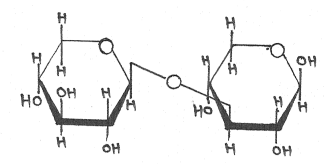


Fig. - 5.

Fig. - 6.

Fig. - 7.

Olicosecharide (3) . a crystalline form having the physical constants identical with those reported for (1 -> 4)-0-7 -0-mylepyranosyl-(1 \Rightarrow 4)-D-mylopyranosa, α -p. 203-05°, $\left[\times \right]_{0}^{22}$ - 46° (in water, C. 1.00%). It was found to be chromotographically pure in solvent F and B. Acid hydrolysis of the eligosecharide yielded only mylose and partial, hydrolysis gave mylose and mylobiose. The Adontity of these oligosepherides was confirmed by ce-chrometography with their authentic cample. The molecular wight was found to be 423, which corresponded to trisaccharide of pentose units. Engymic hydrolymis with emulsia and negative rotation showed that the mylese units were linked through \$ -linkages. The periodete oxidation studies efforded the liberation of 2.2 moles of femic acid and congusption of 5.21 moles of periodate per mole of the trisaccharide. On the besis of above evidences, the elicosaccheride was identified to be $0-\beta$ -0-sylopyranesyl-(1 \Rightarrow 4)-0- β -0 $xylopyranosyl-(1 \rightarrow 4)-0-\beta -0-xylopyranoso.$ (Fig. - 4).

Oligosoccharide (4) a crystalline sugar, m.p. 190°, $\left|\mathcal{L}\right|_{D}^{22} = 30.4^{\circ}$ (in water, C, 2.90°), was found to be chromatographically pure in two solvent systems F and B. The sugar on and bydrolysis yielded only sylose while the molecular weight of the sugar 296 corresponded to a pentose disapeharide. Shaymic hydrolysis with emulsin showed the presence of β -linkage between the two sylose units. The periodete exidation showed the consumption of 3.24 moles of metaperiodete with liberation of 1.15 moles of femmic acid per mole of the sugar. The oligosoccharide is,

therefore identified to be $Q = \beta = 0$ -mylopyrenceyl=(1 \Rightarrow 3)= $Q = \beta = 0$ -mylopyrences. (Fig. = 5). The identity was confirmed by co-chronotography with an authoratic sample.

Cliquesceharide (5), m.p. 186-87°, \mathbb{K}_{0}^{20} = 25° (in water, C, 3.5%), was chromatographically pure in solvent F and 3. Agid hydrolysis showed the presence of mylose only. The molecular weight of the sugar was 290, corresponded to a disappharide of mylose units. The pariodete oxidation showed the liberation of 2.21 moles of formic acid with the consumption of 4.31 moles of metapariodete per mole of oligosaccharide. Hence the oligosaccharide was assigned the structure $\mathbb{G}_{\mathbf{F}}$ $\mathbb{F}_{\mathbf{F}}$ -mylopyranesyl-(1 \Rightarrow 4). Mylopyranese. (Figg- 6). The identity was further confirmed by co-chromatography with an authentic sample.

cliquesceharide (6), [<] 30 + 140 (in water), m.p. 190-910, was shown to be chromatographically pure in solvent 1. On acid hydrolysis revealed the presence of galactose and mylose. The quantitative estimation by the method of hirst and Jones abound the molar ratio to be like between the two sugars in the eliquesceharide. The molecular weight 296, showed it, to be a dissocharide. Foriodate exidation studies afforded liberation of 2.12 moles of fermic acid and consumption of 4.14 moles of periodate per mole of the eliquesceharide. (Fig = 7).

On the basis of the results obtained so far particularity from methylation studies, graded and partial acid hydrolysis, the following valuable information could be derived :

(1) The main chain of the polysaccharide consists of $\beta = (1 \Rightarrow 4)$ and $\beta = (1 \Rightarrow 3)$ linked mylose units.

- (11) All the galactose units are present as terminal groups and linked in the main chain through β =(1 \Rightarrow 4) linkeges.
- (iii) Two mylose units per repeating unit of the polysecheroide are also linked as a side chain and linkages between main chain mylose units and side chain mylose units and side chain mylose units are $\beta = (1 \Rightarrow 3)$.
 - (iv) From the above information, it is also clear that the galectose units in the side chain are linked at the same mylose units in the main chain which linked through $\beta*(1\Rightarrow3)$ linkages in the main chain.

Taking all the experimental evidences into consideration together with the structures of different alignmentation, the following most probable structure has been assigned to the polymercharide from the seeds of Zizyphus jujuba.

$$-\left[4-2yp-\beta-1\right] \rightarrow (4-2y-p-\beta-1)_2 \rightarrow 3-2y-p-\beta-1$$

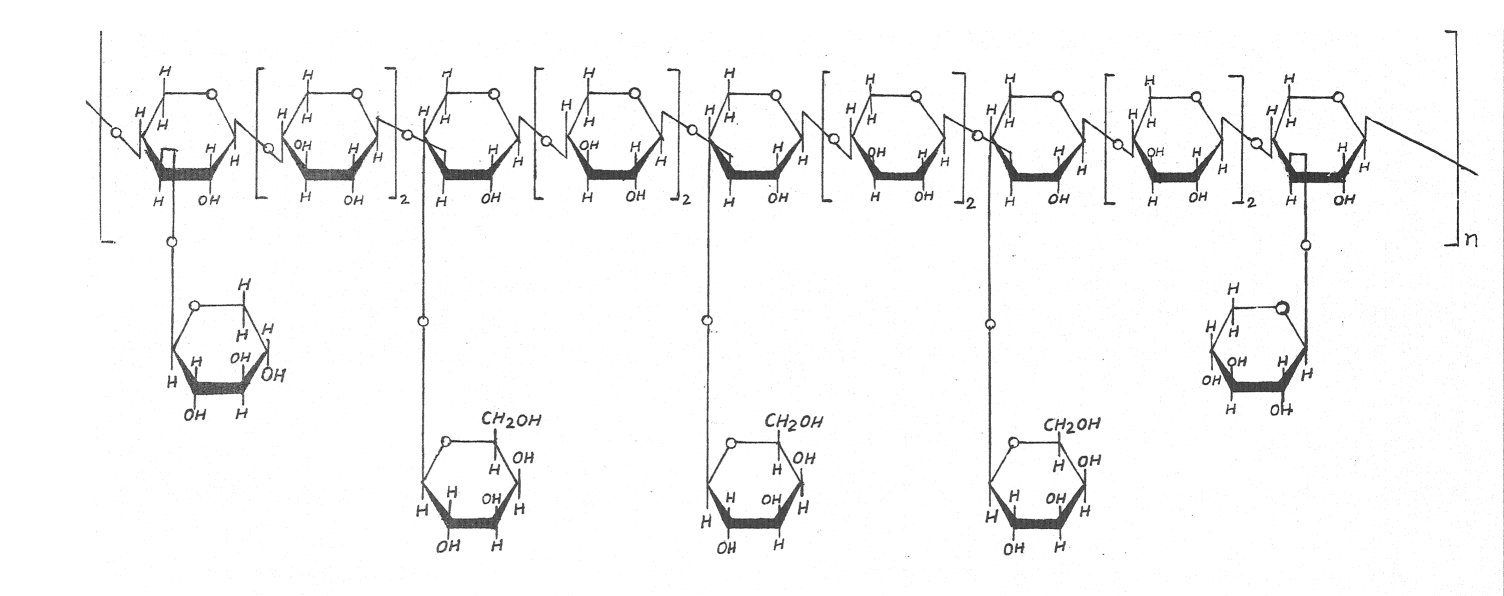
$$-\left[4-2yp-\beta-1\right] \rightarrow (4-2y-p-\beta-1)_2 \rightarrow (4-2y-p-\beta-1)_2$$

Golap a DeGalactopyronose & Nyap a DeNylopyronose.

The above structure contains 10 units of monocarcharides per mapeating unit which fully emplains the formation of alignments. Mides as obtained by partial acid hydrolysis and agrees well with the analytical data of the polysecharide. The dotted and doubly arrowed dotted lines show the probable made of fission of the linkages during the partial acid hydrolysis. The arrowed dotted lines indicated secondary hydrolysis.

The polysoccharide such as described above should consume 18 moles of metaperiodate with the liberation of 5 moles formic acid per repeating unit of 18 auger units. The actual consumpation of periodate 18.21 and liberation of formic acid 5.04 males have been determined for per repeating unit of the polysoccharide, which are in close agreement to the calculated values.

Similar other structures may be possible but they are less probable because the formation of oligosaccherides as obtained in the present case might not be possible.



STRUCTURE OF POLYSACCHARIDE FROM THE SEEDS OF ZIZYPHUS JUJUBA



The state of the s

All evoporation were carried out under reduced pressure at low temperature unless specified otherwise. Residues were dried in vacuum at room temperature over anhydrous calcium chloride. All specific rotations are equilibrium values and all melting points are uncorrected. Paper chromotography was performed at room temperature by descending technique on whotmen No.1 filter paper unless stated otherwise, using following solvent system :

(A)	n-Dutanol - othonol - water	(41115)34
(a)	n-Dutanol - cotic ocid - water	(An 1 alb)
(C)	n-Butanonol- iso-propanol - water	(111623)
(D)	Bonzene - ethanol - water	(169:47:15)
(8)	Butanone - water	
(P)	Sthyl scotate - pyridine - water	(11:4:3)
(G)	Sthyl acetete - pyridice - water	19:1:2)
(24)	n-Outanol - othenol - water	(40:10:19) ³⁴
(1)	n-Sutanol - otherol - water	(5:1:4)

The spots were located by spraying a chromatogram with aniline bydrogen phthelate 43 and heating it at 110-20° for 10-15 minutes. Speckrophotometric determination were carried out by a modification of phonol - subphuric acid method 44. Wheth-Summarson phthelactric colorimeter was used for measureing the absorbance.

LIAS ASSERTION OF THE POLYSHOPHER TOR

The dried and cruehed goods (1.0 %) were extracted successively with petroleum other (60-60°) and othered. The extracted speds were dried and then suspended in distilled water (1 litre) containating 1% acetic ecid. The mixture was stirred mechanically for 8-10 hours to extract the mucilage as such as possible and squeezed out through a muslin cloth. The process was repeated air times when proctically no processive was obtained by adding the extract to an excess of others. The combined extracts was elicated thrice

through a thick cotton pad, placed over a cioth in a Buchner funnel to remove the suspended fine particles. The clear mucilege solution so obtained was added slowly to a large excess of ethenol with constant vigorous stirring when a fibrous colouriese precipitate of the crude polysoccharide was obtained. It was filtered, weshed with ethenol, fellowed by absolute et anol and dried in vacuum at room temperature (31 g s och 3.15%).

11.6 MINIFICATION

The dried crude polysaccharide was dissolved in distilled water (2 litres) containing 15 sectic acid with constant stirring The solution was filtered and was added very alowly to ethenol (8 litres) with constant and vigorous stirring and hept evernight. The precipieted polysaccharide was filtered and the above process was repeated four time, to get a white fibrous muchlage, (25 g ; ach 0.5%).

17-7 HOUNTENETTY OF THE POLYSACTEANTOS

The homogeneity of the polysecharide was checked by the following methods.

11.7.1 (a) Proctional Precipitation

The pure mucilage (4 g) was dissolved in distilled water (500 ml). It was then added slowly to ethanol (500 ml) and the precipieted polysecoheride (Fraction I) was filtered, washed with ethanol followed by absolute ethanol and dried in vacuum. The filtrate was treated with enother 1000 ml of ethanol with stirring and precipitated polysecoheride (Fraction II) was filtered, washed and dried in vacuum. Both the fractions along with the original polysecoheride were bydrolysed separately with Si sulphuric add.

The sugar present in each bydrolyseds wage first identified by

poper chromotography with authorite sugars using solvent (C) and then separated on two sheets of whatman No.1 filter paper using the same solvent. The sugars were eluted with water and estimated quantitatively by periodete emidetion method⁴⁵. The sugars eluted from one sheet were estimated by titration of degmic acid liberated with standard alkali solution whereas the sugars from the other sheet were estimated by the method of converption of periodete. The ratio of Degalactose and Desylose in both fractions was found almost the same (115), indicating the purified polyeaccharide to be homogeneous.

11.7.2 (b) Agetylation and Description

The pure polysaccharide (1.5 g) was mixed theroughly with anhydrous sedium acctate (10 g) and mixture was suspended in acctic enhydride (30 ml). After refluxing over a water-both for 18 hours, the mixture was cooled to room temperature, and poured over crushed ice with constant stirring and then lift overnight. The grayish-white precipiate was filtered, washed with water and dried in vacuum. The diried mass was then dissolved in minimum quantity of acctone and the solution was poured showly in distilled water, where upon a fine fibrous precipiate was obtained. This precipiate was filtered, washed and dried in vacuum 1.12 g.

The dried acetyloted polysoccharide (0.9 g) was dissolved in acetome (32 ml) and 50% potaggion bydrowide solution (32 ml) was added to it. The descetylation was carried out in the usual manner 46 by refluxing the mixture over a water-both for six hours. The viscous solution was poured slowly with etirring into 36 otherelic sold (300 ml) to precipitate the polysoccharide. The precipitate was filtered and was spain precipitated by discolu-

The oxiginal polysaccharide $[\infty]_0^{21} * 91*2^0$ (in water, C. 0.8%) and the polysaccharide obtained after description had almost the identical specific rotations indicating the homogeneity of the polysaccharide.

11.7.3% (c) Zono - Slectrophoresis

A strip support (15 cms x 45 cms) of Whatman No.1 filter paper was marked with a pencil in middle to indicate the starting line. 0.3% solution of polysaccharide (50 ml) was placed on starting line as a compact band. After drying at room temperature the strip was aprayed with borate buffer (pH 9.3) and suspended h horizontally in the electrophoresis tank containing two electrode compartments each having approximately 400 al of borate buffer (pH 9.3). After electrophoresis at 260 V and 12.5 mA for 6.5 hours, the paper strip was dried. It was then cut lengthwise into l on segments, which were numbered to the cathode end. The material from each numbered strip was cluted with water (6 ml) and filtered through glass wool. The filtrate (5 ml) was placed in a hard glass boiling tube with 8.3% aqueous phenol (1 ml). To the tube, concentrated sulphuric acid (15 ml) was added rapidly. The tubes were allowed to cool at room temperature. The absorbance of characteristic yellow orange colour was measured in a Klett-Summerson pnoteelectric colorimeter using filter No.50. A blank was also gun under the same conditions but without polysaccharide.

The meading so obtained were plotted against the segment number counted from the snode end to the cathode end. Only one shapp peak was obtained indicating the polymercharide to be homogeneous.

TABLE .

No.	Klatt reading of alute	Mank Hett Roading	Commocted Klote reading	Abouthman
1	23	22	3.0	0.006
2			1.0	0.002
3			1.0	0.002
4			2.0	0.004
	23		2.0	0.004
6			2.0	0.004
7			3.0	0.006
8	24		3.0	0.006
9			2.0	0.004
10	28	22	1.0	0.002
11			2.0	0.004
12		24	3.0	0.006
13		22	3.0	0.006
14	40	23	17.0	0,034
23	40	22	26.0	
16	39	23	16.0	0.037
17	20		3.0	0.005
	23	21	2.0	0.004
19		2	2.0	0.004
20	24	28	1.0	0.002
57	23	22	1.0	0.002
22	23		2.0	0.004
28				0.006
24		21	3.0	0.006
			2.0	0.004
26		22	3.0	0,006
27		21	1.0	0.002
20	22		1.0	.0.002
20			2.0	0.004
30		2	2.0	0,004

Absorbance was measured on 5 ml portion of coloured solution.

Absorbance = 2_K_ilett_mending.

II.O ASH CONTON

The deted polysaccharide (0.2~g) was ignited in a tilical crucible previously heated to a constant weight. After ignition the crucible was cooled in a deciceator and weighed. From the weight of residue (0.0010~g), the ash content was calculated to be 0.5%.

11.9 PIN'SIGAL AND CHEMICAL EXAMINATION

It was a fibrous white pendered, very light in weight, slowly soluble in water, $[<]_0^{2l} = 91.2^\circ$ (in water, C.0.8 g per 100 ml of solution). For the purpose of optical rotation, the solution was filtered through a sintered funnel to get a clear solution and the amount of polysescharide in the solution was determined colorimetrically. The polysecharide was found to be free of nitrogen, sulphur and halogens. It did not reduce Fehling's solution.

II.10 EXPLEMATION OF PRICE SUGARS

The polysaccharide was examined for free augars by applying three spots of its solution in water on a strip of thatman No.1 filter paper (15 cm x 45 cms). The paper was developed in solvent (A) for 36 hours, dried and cut lengthwise into three strips, each containsing one spot. The three strips were aprayed with three different reagents using naphthoresorainel and trichleroscotic acid (gives colour with hetoes only) on one, aniline hydrogen phthelate on the second and silver nitrate in acetone followed by ethebolic sodium hydroxide. On the third. The first two paper dried in

the oven at 120° and the third was air-dried. None of the strip showed any spot, hence the polysaccharide did not contain any free sugar.

11.11 METHONYL GROUP DETERMINATION

The percentage of methoxyl groups was determined by the method of Belcher, Fildes and Nutten 49 and was found to be 0.74%.

11.12 ACETYL GROUPS DETERMINATION

The method by Belcher and Godbert was followed for the determination of equiple group percentage with and without mucilage. Found acetyl 0.90%.

11.13 URCHAIDE CONTENTS DETERMINATION

The uronide contents were found to be negligible by the semi-micro method of Barker, Foster, Siddiqui and Stacey 51.

11.14 HYDROLYSIS OF FOLYSACCEARIDE AND DETERMINATION OF MONOSACCHARIDES

The purified sucilege (1.2 g) was dispolved in 2N sulphuric acid (100 ml) and was hydrolysed on a water-bath for about 24 hours. The hydrolysets was neutralised with berium carbonate, filtered and concentrated under reduced pressure. The hydrolysets was examined for monosaccharide as described on next page.

IX-14-1 (a) Paper Chromatography

The spots of the hydrolysate were applied on two sheets of whotman No.1 filter paper. The papers were developed separately in solvents (A) and (B) by descending unidimensional technique. The chromatograms were air-dried and sprayed with aniline hydrogen phthelate. On heating them in an even at 120°, each chromatogram showed two spots. The Rg and Rg values of the two spots corresponded to Degalactose and Degyloge as given in the following Table.

1/01/0 - 2

Sugar identified	fould		Acres .	35	
D=Gelactose	0.08	0.07	0.16	0.16	
D-xylose	0.14	0.15	0.27	0.28	

G = 2.3.4.6-Tetra-O-methyl-O-glucose.

The identity of two sugars was further confirmed by cochromatography with authentic samples of the sugars.

II.14.2 (b) Column Chromatography

A portion of hydrolysate was dissolved in a small amount of aqueous methanol (1:1) and adsorbed over a well weahed column of cellulose (1° x 15°). The column was left over-night and the separation was diffected with solvent (A) and several fractions (15 ml) each ware collected. Each fraction was analysed by paper chromatography with authoritic samples of D-galactose and D-sylone

in solvent (8). The fraction 1 = 10 containing same sugar ware combined together and concentrated to give 3-zylose. It was zerozystallised from aqueous methanol, $[K]_{0}^{30}$ + 17.5° (in water ; G, 1.146). The multing point of the sugar was found to be 143-44° The following derivative was propaged.

D-Myloso Phenyl Osezone Derivetive

The oserone of the sugar wee prepared by heating (250 mg) of sugar , 50 mg of phenyl hydrazine hydrochloride and 0.3 g of sodium acetate dissolved in 5 ml of water in a test tube and heated for 30 minutes on a boiling water-bath. Precipitate of the oserone started appearing after 7 minutes. The floogulant precipiate was separated with water, recrystallised from 50% ethenol, m.p. 160-61° resembling to an authoratic.

The fraction 15 = 35 were mixed and concentrated to give D-galactose. It was recrystallised from equations methanol, $\left[\times \right]_{D}^{25}$ + 79.2° (in water, G, 0.5%). The multing point of the sugar were found to be 167° . The following derivatives were prepared.

(1) D-Galagtose Phonyl Hydrazone

Found Beps 153-54⁰ Given (11t.)⁵³

(ii) N-p-Hitrophonyl-D-Galactonylamina

In a microtest-tube were taken galactose (25 mg), p-mitroemiline (25 mg), a drop of glacial acetic acid and 2 drops of methanol : water (6:1 m/v). The mixture was boiled for 8 minutes and
kept evernight in a refrigorator. The crystalline product was
filtered, washed with cold otherol, other and dried in vacuum. It

molted at $218 \text{--}19^{\circ}$ ofter recrystallisation from methanol. Lit. 54 m.p. 219° .

11.14.3 (c) Thin-layer Chromotography

The plates were prepared from slurry of milics gel G in O.1N colution of boxic sold and the spots of hydrolysate along with benzene accetic acid:methanol (1:1:3) and air-dried. These plates were sprayed with amiliae hydrogen phthalete reagent. On heating them at 120° in an oven, two spots corresponding to D-galactose and D-mylose were observed.

II.15. QANTITATIVE ESTIMATION OF MONDSACCHARIDE

The method due to Hirst and Jones 45 was applied for quanti-

The polysaccharide (200 mg) was dissolved in 2N sulphuric acid (20 ml) in a 25 ml round bottom flagk. The flask was then heated for 24 hours on a water-bath. After cooling to room temperature the hydrolysate was diluted to 30 ml and then D-ribese (20 mg) was added to it. The whole solution was shaken well and transferred to a beaker. The flask was washed well with water, and the washings were transferred to the beaker. The solution was netrallised with berium carbonate and filtered. The filtrate and the washing of berium carbonate were concentrated and then made upto 10 ml.

Six sheets (30 x 45 cms) of Whatmen No.1 filter paper wage used as paper chromatograms. Three quide straps (4 x 45 cms) two on either edges and one in centre, ware marked on each paper. A portion of above splution was placed along the starting line.

is one away from the upper edge) of the three sheets, whereas the remaining three sheets were used as blanks. A guide spot was placed in the centre of each guide strip. All the sheets were developed in solvent (C) for 46 hours. After drying the chromatograms, guide strips were cut lengthwise, sprayed with aniline hydrogen phthelate and heated in an oven at 120° to locate the position of sugars. With the help of these guide strips, appropriate sections of uneproyed portion were cut along with the blank strips of same dimensions from the blank chromatograms. Each section (with and without sugar) was cut into small pieces and extracted separately with 10 ml of hot water. The cluted sugars were then exidised with 0.25 H sedium metaperiodate (5 ml). The liberated formic acid was titrated with standard alkali, after destroying the excess of metaperiodate with ethylene glycel (2 ml), using methyl red as indicator. Slank readings were substructed to get the titre values.

Stages	Value used (of alka in ml)	Correspo of sugar	nelog ((45 m	
Galactosa	3.00		0.606	1.200	1.150
Myloso	14.62	20.40	4.501	(,22)	5.812
altone	1.76	2.44	0.542	0.751	0.696

^{*} Strength of sodium hydroxide = 121.8 *

Assuming complete recovery of D-ribose, the above results indicate that in the polysoscheride D-galactose and D-rylose are in the molar ratio of 1:5.

11.16 GRADED INDECLYSIS³⁶ OF THE POLYSACCHARIDE

The polysacchuride (100 mg) was dissolved well in 0.05% sulphuric scid (20 ml) and the hydrolysis was carried out ower a boiling water-bath. The hydrolysates, taken out at various inter-vals, were examined chromatographically without removal of sulphuric scid using solvent (3) for the purpose of irrigation of the paper. Results are given in Table * 4.

(ine (in minutes)		To a come
\$	Gelectose (Feint)	4 (1986)
	Goladtoco + zylogo (Faint)	
	Samo as abovo	
20	Same as above	
25	Same as above	
30	Same as above	Two spots of oligo- sccharids.
60	Sane as above	Two spots of oligo- sechorids.
90	Same as above	Three spots of oligo- scabaride
120	Galactose + mylose	Four spots of oligo- scohoride.
130	Supple as above	Same as above
240	Saao ee above	Same as above
420	Samo as above	Sana as above

During graded hydrolysis of the polysacchamide galectome was found to be liberated first followed by mykeso. The mounts earliest release of Degalectoms and simultaneously of Demykeso

II.16 GRADED HYDROLYSIS³⁶ OF THE POLYSACCHARIDE

The polysaccharide (100 mg) was dissolved well in 0.05% sulphuric edid (20 ml) and the hydrolysis was carried out over a boiling water-bath. The hydrolysates, taken out at various inter-vals, were exemined chromatographically without removal of sulphuric acid using solvent (3) for the purpose of irrigation of the paper. Results are given in Table * 4.

(in minutes)	Sargest Leganitation	No. of other
	Gelectose (Feint)	
10	Goladtose + zpylose (Faint)	
28	Same as aboye	
20	Samo as above	
23	Same as above	
30	Same as above	Two apots of oligo- eccharide.
60	Same as above	Two spots of oligo-
90	Same as above	Three spots of oligo- socharide
120	Galactose + mylose	Four spots of oligo- eccharide.
130	Samo as above	Samo as abovo
240	Samo es above	Same as above
420	Samo as above	Same as above

During graded hydrolysis of the polysochamide galactose was found to be liberated first followed by mydess. The mounts earliest release of Degalactose and simultaneously of Demylose

(faint) leads to the conclusion that Degalectose are present as terminal groups and some units of Degalectose are also present as terminal groups instead of main chain of the polyseccharide. As galectose is liberated explica than mylose, this is most probably attached to the main chain by more easily hydrolyseble linkages.

11.17 METRYLATION OF POLYSACCHARIDE

The polysaccharide was methylated first by the method of Parikh, Ingle and Shide 57 followed by Furdie's method 56.

The polyearcharide (8.0 g) dissolved in minimum amount of water and then taken in a conical flask fitted with 8*24 joint.

Dimethyl sulphate (40 ml) and 40% sodium hydroxide (80 ml) were added dropwise with constant stirring with magnetic stirrer. The temporature was maintained between 40*50°. After repetition of the above procedure, the solution was concentrated under reduced pressure and filtered to remove the sodium sulphate. The filtrate was again concentrated to a thick syrup and dissolved in accorne. This was then methylated by repeating the above procedure thrice. The finally concentrated solution was extracted thoroughly with chloroform. The extracts were dried over anhydrous sodium sulphate and the solvent distilled off under reduced pressure. The partly methylated product was brownish mass. (6.92 g). *OCH₃ : 35.5%.

The partly methylated polysaccharide was further methylated by Furdie's method. The partly methylated polysaccharide (6.5 g) was dissolved in methanol (36 ml) in a conical flock fitted with three method multiple adapter. The temperature was maintained at

40-50° by placing the conical flash, fitted with air-condenser having fused GaGl₂ -tubes in a trough containing water over the magnetic stirrer. Methyl iodide (9 g) and silver exide (6 g) were added with continuous stirring in several equal instalments, each after helf an hour interval. After the final addition the reaction mixture was heated for four hours on a water-both under reflux and then filtered after cooling the contents. The silver salts were exhaustively extracted with chloroform under reflux. The combinedfiltrate and extracts were eveporated under reduced pressure and the resulting syrup was remethylated thrice under the same conditions. The fully methylated polysoccharide was obtained as a deep becom coloured product. (5-1 g) +0GH₃, 44-66 a [c] 21 - 36° (in chloroform, G, 1-0 g per 100 ml of solution).

II.18 IMDROLYSIS OF THE METRYLATED POLYSACHARIDE AND IDENTIFICATION OF METRYLATED SUGARS

The hydrolysis of methylated polysaccharide was carried by slight modification of method due to Souveng etcal³⁹. The methylated polysaccharide (100 mg) was dissolved in 85% formic scid (20 ml) and solution was refluxed for 4 hours on a water-bath. The solution was then cooled and concentrated under reduced pressure and traces of formic scid were removed under vacuum. It was dissolved in 0.23 % sulphuric scid (10 ml) and the hydrolysis was carried out for 16 hours on a water-bath. The hydrolysis was carried out for 16 hours on a water-bath. The hydrolysate was & cooled, neutralised with barium carbonate and filtered. The residue was washed with water followed by ethanol. The combined solutions wase concentrated under reduced pressure to light brown symp. The methylated sugars wase separated on Whatman No.1 filter

paper using solvent (A). The chromotograms showed four spots, after spraying with amiliae hydrogen phthalate and drying at 120°. The R_{DIG} (THG = 2.3.4.6-totra-0-methyl-0-glucose) value of each methyloted sugar was calculated in solvent (A) and R_g value was calculated in solvent (A) and R_g value was calculated in solvent (I). These values were compared with that given in literature as shown in the following table.

Methyloted sugare identified	TAG Lound	TMG 34,61
2-0-Methyl=0-xyloss	0.30	0.38
2,3-Di-0-mothyl-D-mylose	0.76	0.74
2,3,4-Tgl-O-mothyl-Demyloge	0.92	0.94
2,3,4,6-Tetra-O-methyl-D-galactose	0.90	0.88

11.19 QUANTITATIVE ESTIMATION OF METHYLATED SUGARS

The methylated polysaccharide (300 mg) was hydrolysed as described above. After hydrolysis, glucese (60 mg) was added to hydrolysate. It was then neutralised with barium carbonate and filtered. The residue was washed with ethanol. The filtrate and washings were concentrated under reduced pressure to a syrup. A portion of the syrup was dissolved in acetone and applied on three sheets (A, B, and C) of whether No.1filter paper. Each having three guide strips. The papers were irrigated with solvent (D) along with three blank sheets. After development of chromatograms and locating the sugars on guide strips, appropriate sections, containsing sugars were cut from the unsprayed protion of the chromatograms. The sugars were cluted with 10 ml of water.

The mothylated sugars were estimated by alkaline hypolodite method⁶⁰. The eluted portions were taken in 50 ml consical flashs separately provided with ground glass joint stoppers and a solution (2 ml) containing 0.2% sodium bicarbonate and 0.2% sodium carbonate was added Solution of iodine (0.1%, 2 ml) was then added to the reaction mixture and the flask was stoppered. The experiments as corresponding blank elustes were also carried out in the same way. After three hours, the reaction mixture was acidified cautiously with 2 N sulphuric acid and 15% potessium iodide solution (2 ml) was then added to it. The liberated iodine was titrated against 0.01N sedium thiosulphate solution using starch as indicator. The results are given in Table * 6.

	ection & Sugar		or 0.0 ad (10			oneling er (£n d	esoune 0)
A	2,-0-mothyl-0-zyloso	2.30	3.13	2.74	1.722	2,321	2.000
	2,3-01-0-methyl-0- xylose	3.46	4,60	4.00	2.760	3.744	3.200
0	2,3,4-Tri-O-mothyl D-mylogo	0.78	1.00	0.92	0.678	0.922	0.800
0	2,3,4,6-letra-0- methyl-0-xylose	0.96		1.10	1.046	1.373	1.199
	Glucoso		1.06	2.62	1.243	1.674	1.430

The above results correspond to an average molar ratio between A : B : C : D as 2.5 : 4 : 1 : 1.5 or 5 : B : 2 : 3. The methylated sugars were calculated as the methyl ethers of anhydromethose and anhydromentose units i.e. $C_0 I_{10} O_4$. $C_2 I_{12} O_4$ and $C_3 I_{14} O_4$ for mone , dim and trimO-mathylaD-maylose respectively and

 $^{\rm G}10^{\rm H}18^{\rm G}_{\rm S}$ for tetre=0-methyl=0-galactose. An average recovery of the methylated polysaccharide was found to be 99.60% assuming 100% recovery of D-glucose.

11.20 CHARACTERISATION OF METHYLATED SUGARS

The methylated polyencheride was hydrolysed according to the method of Garege and Lindbarg 62. Nothylated polyencheride (4.0 g) was dissolved in 72% sulphuric acid (50 ml). The solution was kept for one hour at room temperature (25°) and then diluted to 200 ml. Further hydrolysis was carried out by heating for 4 hours on a water-bath. The solution was cooled neutralised with barium carbonate and filtered. The residue was mashed with water followed by ethanol. The solutions were concentrated to a syrup under reduced pressure.

The mixture, containing different methylated augers, was resolved into five fractions on whetman No.3 filter paper using solvent (D). Strips, containing different individual methylated sugars, were eluted with water. The eluates were concentrated separately under reduced pressure and marked as fractions, I, II, III, and V.

II.20.1 Proction I

Solid, $R_{\rm DIG}$ in solvent (A), 0.39, (No. 18.96%, calculated for mono methyl pentose, $G_{\rm c}H_{12}G_{\rm p}$ CMs , 18.90%, m.p. 130-32° [K] $_{\rm D}^{\rm 25}$ - 24° (in water, C, 25). Lit. Supp. 133-37° [K] $_{\rm D}^{\rm 25}$ - 23 ->+35° (in water). Lit. Supp. 133-33°, [K] $_{\rm D}^{\rm 25}$ - 24 ->+36° (in water). It formed 2-0-methyl-1>-mylose and lide on treatment with athenolic

and line, m.p., $123-24^\circ$, $[\propto]_0^{25}$ = 213° (in othyl scatate solution, C. 0.8%). Lit. 63 , m.p. $123-26^\circ$, $[\propto]_0$ + 214° (in othyl scatate).

On acceptation of sugar with anhydrous sodium accetate and accetic anhydride a groyich white precipiate was obtained. The dried mass dissolved in minimum quantity of accetone and the solution was poured slowly in distilled water, where upon a white crystalline compound 2-0-mathyl-D-mylose; 3,4-discotate, m.p. 73-77° $\cdot \left[\times \right] = 38.5°$ (in chloroform, C, 2.5%). Lit. 64 m.p. 76-79° $\cdot \left[\times \right] = 38.5°$ (in chloroform).

11.20.2 Proction II

Syrup, R_{TMG} in solvent (A), 0.76 , found (No. 34.8% dimethyleylose, $G_7H_{14}G_5$, requires with, 34.8%. The optical rotation of the augus [K] $_0^{20}$ + 22.2° (in mater, G, 4.3%), Lit. 67 , [K] $_0^{18}$ + 23°.

The anilide of the sugar prepared by the method of Hampton to dry syrup (200 mg) were refluxed for aix hours with 1.5 ml of freshly distilled dry aniline dissolved in 10 ml of absolute ethenol. The ethenol was distilled off and the bulk of the aniline was removed under high vacuum, (5-6 mm of mercury) at 65-70° (buth temperature). The syrup mass was kept in the referigerator for 72 hours, when tiny (respectals (plotes) were observed. The adhering aniline was removed by the addition of dry ether, and the crude crystals (light brown) were filtered out, weshed with other and dried. (yield 40 mg), m.p. 138°. Lit. 67, m.p., for 2,3edi-O-mothyl-D-mylopyranosyl smilide is 145° and optical

motation $[\[mathcal{C}]_0^{21}$ + 192.3° (in othyl acotate, C. 0.223%). Lit.⁶⁷ $[\[mathcal{C}]_0^{14}$ + 190° (in othyl acotate) and in Lit.⁶⁹ $[\[mathcal{C}]_0^{1}$ + 185° (in othyl acotate).

The methoxyl consent of the 2.3 - di-0 - methyl-0-xylopyreness anilide (recrystallised was found to be <math>25.2% calculated for $C_{13} H_{19} O_4 H_8$ (No. 24.0%). The sugar present in this fraction was identified as 2.3 - di+0 - methyl+0-xylose.

11.20.3 Proction 111

Syrup, it could not be recrystallised. The $R_{\rm DIG}$ in solvent (A) 0.92, optical rotation of sugar was found to be [4] $_0^{18}$ + $_18.2^{\circ}$ (in water, G, 0.39%), Lit. $_0^{18}$ is [4] $_0^{18}$ + $_20.3^{\circ}$, one found , $_35.12\%$, calculated for $_3H_{16}O_5$ is $_35.35\%$.

The emilide of the sugar was propored by refluxing the dry syrup (38 mg) with freshly distilled dry emiline (120 mg) for three hours in a water-bath (85-95°) in absolute ethenolic solution (5 ml). Sthemol was distilled off and the whole thing was kept in the refrigerator for seven days. The 2,3,4-tri-0-mathyle-b-sylopyrenosyl emilide failed to crystallise. It cans out as a white pro powder by the addition of 3-4 drops of drys scatome. The precipitate was filtered out and dried, (yield 12 mg). The m.p. of powder was found to be 94.96°, $[\infty]_B^{22}$, 83° (in ethenol, C, 26). Lit. 67 m.p. 120°, $[\infty]_D^{-}$ = 84 $^{-1}$ + 47° and Lit. 70 m.p. 91°. The mathomyl value of the derived emiliate was found to be 33.96 ($C_{14}^{12}C_{11}^{12}C_{11}^{13}$ requires, $C_{12}^{12}C_{11}^{13}C_{11}^{13}$ requires, $C_{12}^{13}C_{11}^{13}C_{11}^{13}$ requires, $C_{12}^{13}C_{11}^{13}C_{11}^{13}$ requires, $C_{12}^{13}C_{11}^{13}C_{11}^{13}$ requires, $C_{12}^{13}C_{11}^{13}C_{11}^{13}$ requires, $C_{12}^{13}C_{11}^{13}C_{11}^{13}$ requires, $C_{12}^{13}C_{11}^{13}C_{11}^{13}$ requires, $C_{12}^{13}C_{11}^{13}C_{11}^{13}$

The sugge in this fraction was thus identified as 2,3,4+
twi-0-mathyleDemyloop.

II.2D.4 Proction IV

A solid, \$\mathbb{B}_{BMG}\$ in solvent (A), 0.90, found the, \$1.66.

Calculated for tetrasethyl hences, the, \$2.46, \$\sqrt{2}\$ + 126

(in water, \$C\$, 0.46). Lit.\$^{71.72.73}\$ for \$2.3.4.6**tetra*0**enthyl**

galactors, \$\mathbb{C}_{D}^{16} \times 142^{\circ} \times 117^{\circ}\$ (equil.) in water, \$\mathcal{C}_{L}\$ 1.15),

m.p. 70=72^{\circ}. It gave a red colour with premisidine hydrochloride aproy in butanol and a brownish red colour with aniline hydrogen phthalate. Its treatment with alcoholic aniline gave \$2.3.4.6**

tetra*0**mathyl**i**phanyl \$\mathref{D}\$-galacosylamine, \$m.p.\$ 180**90^{\circ}\$, \$\mathref{C}_{D}^{\circ}\$**

80^{\circ}\$ (in acetone, \$\mathref{C}_{L}\$ 1.06), Lit.\$^{71}\$ m.p. 193**94^{\circ}\$, Lit.\$^{74}\$ m.p. 192^{\circ}\$,

II.-21 PERICOATE ONDATECH OF DIE FOLYSACCHARIDE

II-21-1 (a) ilberation of formic acid 30 and estimation of end oroug

The polysocharide (500 mg) was dissolved in water (5 ml) and in this solution, potessium chloride (0.5 g) and 0.23M sodium metaperiodate (60 ml) ware added. The volume was made upto 140 ml with water. In a blank apperiment, potessium chloride (0.5 g) and (0.23M) sodium metaperiodate (60 ml) ware diluted to 140 ml with distilled water. The oxidation was carried out in dark at room temperature. 5 ml. of aliquots were drawn at various intervals along with blank and emess of metaperiodate was reduced with 2 ml of ethylene glycol. The liberated formic acid was titrated against W/110 sodium hydroxide using methyl red as indicator. Results are given in Table * 7.

The data shows that 0.2066 male of formic acid was liberated (72 hours) per 100 g of polysaccharide. The ascent of formic
acid liberated (72 hours) corresponds to 20.33% of anhydrohenese
units present as end groups. The titre value of sikeli at 48.60,
and 72 hours indicated that one male (of formic acid was liberated
per 531.0 g. 491.1 g and 464.03 g of the polysaccharide respectively.

in loure)	Roading wi blanks (in ml)	th Volume of elkeli uged (in mi)	formic seid linerated	lotal formic sold liberated (in mg)	
	0.0	2.06	1.196		
	0.0	3.06	1.279	35.612	
24	0.0	3-26	1.360	30.164	
36	0.0	3.48	1.450	40.600	
40	0.0	3.70	1.347	43.316	
60	0.0	4.00	1.073	46.844	
72	0.0	4.0	1.697	67.516	
	0.0	4.00	1.697	47.516	
96	0.0	4.00	1.697	47.516	

After 72 hours, 25 ml portion of reaction minture was taken out, acidified with 2% sulphuric acid (5 ml) and then 10% potassium iodide (4 ml) was added to it. The liberated iodine was titrated immediately against 1% sodium thiosulphate solution without using starch as indicator till the solution became colour-less. The solution was concentrated to 10 ml to which 2% sulphuric acid (10 ml) was added and the hydrolysis was carried out for 16 hours on a water-both. The hydrolysis was carried with

berium carbonete, filtered and the filtrate was concentrated to a syrup under reduced pressure. The syrup was examined by paper chromatography using different solvents the chromatogram revealed the presence of nylose only, galactogo found to be absent completely.

II.21.2 (b) Consumption of Metaperiodate 76

The polysaccharide (250 mg) was dissolved in water (70 ml) to which 0.25 M sodium metaperiodate (40 ml) was added and the total volume was made upto 120 ml with water. A blank was also prepared with 0.25 M sodium metaperiodate (40 ml) diluted to 120 ml with water. The periodate exidation was carried out at room temperature. 2.0 ml aliquots were withdrawn from the reaction mixture and blank at various intervals and to them 20% potassium iedide solution (2 ml) was added followed by addition of 0.5M sulphuric acid (3 ml). The liberated iedine was titrated immediately against 0.0404M sodium thiosulphate solution using starch as indicator. The reading with the polysaccharide were substracted from the corresponding readings of control experiment to get the titre values. The results are given in Table 8.

TADLR - 0

1100	Volume of hypo Corresponding used smount of		Total periodate consumed		
hours	(in al).	periodata consumed (in mg)			
(1)	1.02	4,409	264.55		
16	1.10	4.759	295.30		
24	1.10	5.100	306,00		
36		5.446	326.60		
48	1.3	9.702	347.55		
•	1-0	4-442	300-11		

TABLE - 8 (Continued)

(in hours)	Volume of used (in ml)	hypo Corresponding amount of periodate consumed (in mg)	lotal paraodate tongund
72	1.46		378.07
34	1.54	6,667	399-42
96	1.54	6.657	399.42

pends to the consumption of 0.7466 moles periodate per 100 g of polysaccharide. After 84 hours periodate oxidized solution (10 ml) was hydrolysed with 28 sulphuric seld (page 37). The hydrolysete was examined chromatographically for the presence of Degalectose and Degalectose. The chromatographically for the absence of both the sugars.

11.22 PARTIAL ACID HYDROLYSIS OF POLYSACCHARIDS

The polysaccharide (6 g) was suspended in water (500 ml) in a three necked flask and was dissolved stirring mechanically. The hydrolysis was carried for four hours at 80° by adding 0.2% hydrochloric acid (5 ml) and the solution was stirred throughout the process. The contents, after cooling down at room temperature were poured in ethanol (2 litres) to precipitate the degraded polysaccharide. The precipitate was filtered and washed well with ethanol. The filtrete and washings were neutralised with silver carbonate with stirring. The precipitate was filtered, washed with water and the combined solutions were concentrated under reduced pressure to a syrup.

11.22.1 Examination of the Precipitate

The precipiate was hydrolysed with 2% sulphuric acid for 18 hours, over a water-bath. The hydrolysate was cooled, neutro-lised with berium carbonate and filtered. The filtrate and washings were concentrated and examined chromatographically over Whatman No.1 filter paper using solvents (A) and (C). The chromatograms showed two spots corresponding to Rg values of D-galactose and D-mylose, which was confirmed by co-chromatography with their authentic samples. Due to small amount of precipiate, further studies were not possible.

11.22.2 Unamination of the Hydrolynate

The hydrolyeate was examined paper chromatographically using solvents (A), (B), (C) and (G). The chromatograms showed seven spots on spraying with aniline hydrogen phthalate and drying at 120°, indicating the prosence of seven sugars.

11.22.3 Separation of Oligomagharides

The syrup was dissolved in minimum quantity of water and applied on twenty sheets of Whetman No.3 paper as long thin band, three inches below the upper and and one inch away from the outer edges. Each paper has three guide strips, two on outer edges and one in centre. After developing the paper on solvent (3), for sixty hours, they were dried. The guide strips were cut from the chromatograms, aprayed with aniline hydrogen phthalate and dried at 120° with the help of the guide strips appropriate sections were cut from the unepreyed parties of the chromatograms and sugars

were eluted with water. In all, soven fractions were obtained.

Mi-22.4 Examination of Fraction I and Identification of Mylotetrose (3 2 - β -Mylobiosylaylothose)

 $R_{\rm g}$ values were 0.62 and 0.06 in solvents (F) and (S) magportively, and mylotetrose values were 2.3 and 2.2 in solvent F and
(B) mespectively (Page 31). \sim \sim \sim \sim \sim \sim \sim 37.80° (in water, C, 3%).

The sugar was hydrolysed with 2N sulphuric acid, neutralised with barium carbonate and filtered. The filtrate was concentrated and examined by paper chromatography using solvents (A) and (B). The chromatograms showed only one spot corresponding to Bg value of Dawylose. Thus sugar consist of only mylose units. Nelecular weight of the sugar was determined by hypoledite method of, and was found to be 550.3, which corresponded to a tetrasccharide of peatoses. Calculated molecular weigh for G20H34017. 546.

Partial acid hydrolysis of tetrosaccharide gave two trisaccharides and two disaccharides which were identified by their R_N values and co-chromatography with their authentic samples. These fractions were trisaccharides of $0-\beta$ -D-mylopyranosyle $(1\rightarrow 4)-0-\beta$ -D-mylopyranosyle $(1\rightarrow 4)-0-\beta$ -D-mylopyranosyle $(1\rightarrow 4)-0-\beta$ -D-mylopyranosyle $(1\rightarrow 3)-0-\beta$ -D-mylopyranosyle $(1\rightarrow 4)-0-\gamma$ -No-mylopyranosyle $(1\rightarrow 4)-0-\gamma$ -mylopyranosyle $(1\rightarrow 4)-0-\beta$ -D-mylopyranose, and $(1\rightarrow 4)-0-\beta$ -D-mylopyranose.

The presence of one (1 -> 3) limitage between two mylese we units in the tetrosecoharide was further confirmed by periodete endiation which showed the consumption of 5.2 moles of netaporiodate.

Sedete with the liberation of 2.16 moles of formic acid per mole of oligosaccharides. The oligosaccharide was completely hydrolysed with emulsin, suggesting | 3 - glycosidic linkages in the oligosaccharide molecule.

All the above results indicates that the oligometheride is 0- **-xylopyranosyl*(1 \rightarrow 4)**-0**-\begin{align*} 3 \text{**-0**-\beta} = 0 \text{**-xylopyranosyl**(1 \rightarrow 3)**-0**-\beta} = 0 \text{**-xylopyranosyl**-(1 \rightarrow 3)**-\text{**-xylopyranosyl**-(1 \rightarrow 3)**-\text{**-xylopyranosyl**-(1 \rightarrow 3)**-\text{**-xylopyranosyl**-(1 \rightarrow 4)**-\text{**-xylopyranosyl**-(1 \rightarrow 4)**-\text{

The identification of sugar is well supported by its cometents found and reported in literature shown in the following Table = 9.

Constants	Found	Reported	Reference
W.			(77)
optical motation [× 21 - 57.8°	[4] = 56.7%	L ^O (77)
R in solvent (F)	0.62	0.63	(39,77)
R _k Xylotetraces in solvent (3)	2.2	2.1	(39,77)

II.22.5 Examination of Fraction II and Identification of 32-B-xylosylsylobiose

This fraction was crystallised from ethanol, $m_*\rho_*$ 222° and $[\prec]_0^{21}$ = 51°kn water, G. 2.9%). Nylotrices values were 1.38

and 1.41 in solvents (F) and (S). $R_{\rm g}$ values in solvents (F) and (S) were found 0.70 and 0.22 sespectively.

The complete eqid hydrolysis with 2% sulphuric eqid, subsequent neutralisation with barkus carbonate and examinationy paper chromatography indicated the presence of mylose only, which was further confirmed by co-chromatography with an authentic sample. The molecular weight of the sugar was found to be 420 by hypodedite method which corresponded to triseccharide of pantess units, molecular weight calculated for $C_{15} i_{125} O_{13}$, 414.

Partial hydrolysis of trisaccharide with 0.3N hydrochloric acid 100° for 30 minutes gave sylose, sylobiose, and rhodysease biose. Periodete emidation studies revealed that one make of the eligosaccharide consumed 4.3 makes of metapariodete and 2.1 makes of formic acid liberated. It also confirmed the presence of 1 = 3 linkage between two sylose units in the oligosaccharide molecule.

The sugar was completely hydrolysed with emulsia, suggesting the presence of β -linkage. From the above observations, the sugar was identified to be 0- β -D-mylopyranosyl- $(1 \Rightarrow 3)$ -0- β -D-mylopyranosyl- $(1 \Rightarrow 3)$ -0- β -D-mylopyranose 1.0. 3^2 - β -mylopylaylobiose. The constants of sugar are given below in Table - 10.

Constants	Pound	Papartod Ba	forences
Coptical rotation € 21	222° - 51° [4] ²² -	225° -> -472.1°	(77) (77)
Bayloblose is solventa (F) & (B)	1.39 & 1.41	1.36 & 1.43	(77)
	0.70 & 0.22	0.72 & 0.20	(22)

11.22.6 Exemination of Fraction III and Identification of Mylotricse

 $R_{\rm g}$ values were 0.36 and 0.10 in solvents (F) and (S) respectively. The sugar was recrystallised from 90% ethanol, m.p. $203-05^{\circ}$. 1000° and 10000° and 10000° and 10000° and 10000° and 10000° and 10000°

Acid hydrolysis with 2% sulphuric acid followed by neutrelisation with barium carbonate, and paper chromatographic exeminetion showed the presence of mylose units only. The molecular weight
was found to be 423 by hypoiodite method which corresponded to
a trisaccharide of pentose units, molecular weight calculated for

Clophicola, 414. Feriodate ordetion of sugar revealed the consumption of 5.21 moles of sodium metaperiodate, liberating 2.2 moles
of oligosecharide.

Partial acid hydrolysis with 0.5% hydrochloric acid 100° for 15 minutes resulted in formation of mylese and mylobiose, which were identified by co-chromatography with an authentic sample.

Constan			Roportod	
Q-P-		233-050	203-06°	(77)
Optical	rotation	W] * 40°		(78)
	301vents			
(8		0.36	0.35	(39,77)
and (8)		0.10	0.09	(35,77)

IL-22.7 Samming tion of Exaction IV , and Identification of Ehodymonobiose

Acid hydrolysis of the suger with 24 sulphuris acid and neutralisation of the hydrolysate with barium carbonate followed by paper chromatographic analysis in solvent (C), reveals the presence of ryless only. The molecular weight was determined by hypoiodite method 60 , 296, molecular weight calculated for mylebiose, $G_{10}H_{10}G_{9}$, 282.

The periodate exidetion studies showed the conumption of 3.24 moles of matepariodate with liberation of 1.10 moles of founds ocid. The sugar was completely hydrolysed with smulsin, showing the presence of β -linkage. Its identity was further confirmed by proparing its phenylosumons derivative, n.p. 196-96°, $[e]_{\beta}^{(2)}$ + 0° (in pyridine, G. 25). And calculated for $G_{2}^{(1)} \circ G^{(2)}$.

N.12-18, found 12-30%.

Constante of sugar were compared with those reported in literature as shown in Table - 12.

Sugar or derivative	Constants	found	Reported	References
Rhodymenabiese	0.0	390°	192-98°	(79)
	Rylobicee in solvent (B)	1.99	1.07	(80)
•d0••	optical [2]	22-20-4	[2]	p.6° (80)
3-0-6-0-xylopyme- nosyl-0-xylose- phonyl osazone	m.p.	196-98 ⁰	194-96°	(79)
*do**	ptical rotation	[권 ²² + 49°	MD * 47°	(79)

II.22.8 Examination of Fraction IV and Identification of Mylobiose

The fraction was recrystallised from aqueous ethanol, m.p. 183-84°, [<] $_{D}^{20}$ = 25° (in water, C, 3.5%). R, values were in solvents (3) and (F) 0.32 and 0.85 respectively.

Hydrolysis of the sugar with 2% sulphuric acid and neutrelisation of the hydrolysate with berium carbonate followed by ppear chromatography in solvent (G), revealed the presence of mylese only which was further confirmed by co-chromatography with authentic sample. The molecular weight of the sugar was 298, calculated for ${\rm C_{10}H_{18}O_{9}}$, 282.

The periodate exidation of sugar consumed 4.31 moles of metaperiodate liberation with 2.21 moles of formic acid indicating the $(1 \Rightarrow 4)$ linkage between mylose unit. The polyseocharide completely hydrolysed with emulsin indicating the β -linkage between two units.

Thus the oligosaccharide is a disaccharide composed of D-nylose linked through β -glycoside bond. The sugar was identified $4\text{-}0\text{-}\beta$ -D-nylopyranosyl-D-nylose, which was confirmed by preparing the phenyl esszone derivative, m.p. 204° and $[\ensuremath{\ensuremath{\bigcirc}}]^{25}_0$ - 51.8° (in pyridinesethanol).

The constants of sugar are given in Table - 13.

TANK 13

Sugar or derivativa	Constants	Found	Bapartod	802.
Xylobiose	m.p. Optical rotation	183-85° 20 • 20°区30	185°,187°	(81.)& (73.) (81.)
		20.	32 ² ->	(82,87)
edo.e	a _x in solvent (a)	0.32	0.33	(81)
phenyl osazone	a.p.	204°	2030	(ex)
***	Optical rotation	5- 51-8° [4]D	- 30°	(8T)

II.22.9 Examination of Praction EVIand Identification of O- P-D-Galactopyranopyl-(1 → 4)-O- β-D-gylopyranops 60,66

Syrup, having optical rotation, [30 + 140 (in water).

Acid hydrolysis with 2% sulphumic scid and neutralisation of the hydrolysate with barium carbonate, followed by paper chromatography, mevealed the presence of D-galactose and D-mylose. The quantitative estimation by the method of Hürst and Jones 45 showed the molar ratio to be 1:1 between the two sugars in the eligosaccharide.

Periodete oxidetion studies showed the consumption of 4.35 moles of periodete and liberated 2.1 moles of formic ecid.

Sethylation of the disappharide followed by acid hydrolysis of the fully methylated derivative afforded 2.3.4.6—tetra—O-methylated derivative afforded 2.3.4.6—tetra—O-methylated begalactors and 3.4—dis-Q-methylated in equal proportions. The polysaccharide was completely hydrolysed with equisin indicating the β -linkage between the two units.

These results ; proved that oligosaccheride was 2-0-0galactopyranosyl-D-mylopyranose.

11.22.10 Ememination of Praction VII and Identification of Describes

The R_g value in ware in solvent (B), 0.28 and R_G value in solvent (A), 0.15, m.p. $143 = 44^{\circ}$, [<] 30 + 17.5° (in water, G, 1.145). The sugar was identified to be D-myloss by co-chromatography with an authentic sample.

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A NEW WATER SCHUELE POLYGACCHARIDE FROM

Masgous Pango.

the seeds of <u>Phosolus supro</u> bolomics to the family Leguminosae.

The plant <u>theseeles munon</u>, ideas, is commonly known as the Stone longer and trailing, who has plant height with raddish trown pubescence, which gives the falliage a lighter tint; leaves large; the pade are nearly exact, weary height and seeds are larger and longer than those of sung and usuamally dark brown and sametimes of a dull geomish.

Und is cultivated in the Uppper Gengetic Plain, especially in the Mearut and Robilkhand Division, and some part of Bundelkhand Region.

Seeds used as dist in fover and to strengthen the eye2.

The work done in the past years on this genus was surveyed and the details of it are given in the tabular form on the next page.

		Plant specios	Constituents		
	Phagoolug	acconitrifoliga	Grystellino globulin	Soade	(1997)3
	Phosoolus	engularie	Keenpferol- robinoble-7- rhennople	Leaves	•
	Phageolug	tellobetus	Keenpferel- robinobie-7- rhannogide	Loaves	
4.	Phoseolus	cocciocous	Starch, emyloge 2.7- emylopectic	Rooto	(1963) ₂
			Uridine di- phosphate-Wh acetyl glucose mine and wridin di-phosphate gr uronic said		(1997)6
6.	Phopoolus		Pivo Cligo-		(1963)
	Phecoolus	lunatus	HCN producing coopounds	Sodo	(1915) ¹¹ (1921) ¹² 13,14,1 ⁵ ,
	Resolus		The proteins and charac- terisation of protein.		(1922)16
9.	Phoenolus	lunetus		30000	(192 5)17
	Photockus	aultiflorus	Chonical invostigations on enzyme, oil polysterol etc.		
4	Aspellus	coltificos	Lipid content		(1200) ¹⁹

(Containing)

		Float species	Constituenta	2024	Veloato to
	Phosoolus	coltifleres	Acyl bydroloso	100000	(1999)20
	Megeolus	aultiflorus	ino giberello ino liko compoundo	Loswoo	(1968) ²¹
	Phoseolus	radiotus	Phosphoryless & Geonzyss	Leaves	(1952)2
	Phesoclus	rodiatus	Phosphoglueo	Soods	(1954) ²³
	Phesoclus		Whosphoply- caric edid and 2-phospho- glyconic edid		(1994)
	Massolus		desential saine acids; leucine, leo- leucine, voline, histidine, lysine & typine & t		
10.	Phesoclus	radiabus	Mosture, ash		(1959)26
19.	Massalus	rodiatus	Alkaline & glycerophoe- photos	Soods	(1960) ²⁷
20.	Phasoolus		Gluceso, galactoco, fructoco, reffinoco, stachyono & werboncoco.	Soods and outer sood cant	

		Plant species	Constituents Part	
	Phagoolus	rodictus.	«Closulto & β	(1979)29
***	Phaseolus	wigasio	2-Phosphogly- Leaves colate phos- phohydrolase	(1979)30
20.	Maggolug	vulgorie	Stachyone Seeds	
24.	Phaseolus	vilgaris	L=(*)- pibecolic acid	(1954)32
25.	Phagoolug	vulgarie	Nelecie edd Seeds	(1960) ³³
26.	Masoolug	wilghels	Phagelic agid Soeds	(1960)34
.7.	Thesoolus	wilgaris	Vecilin like, Seeds loguinin like protein	(1979) ³⁵
	Hasoolus	vulgazis	β · Program	(1964)36
.Ve	haseolus	vulgazio	Anino ocide	(1966)37
0.	Phaseolus	vulgazie	2-C-cothyl crythronic ocid	(2979)38
1.			pentothenic ecid & gluconic ecid	
1.	Phasoclus .	wigazio	Aeid phosphoto	(1979) ³⁹
2.	Pheeoclus	wlgeria	Six anthre- Seed quinones coets	(1966)40
)3.	hassolus	wilgesis	Storolic coopds. Coty-	
4.	Phospolus	volgaris	Carbohydrates «do.»	(1831),4
		migazio	Gila Cotyled	ono (1932) ⁴

Different perts of this genus have been investigated for different plant products as has already been described in literature, but no neutral polycecharide has been mentioned uptill now. Therefore an attempt has been made for isolation and structure elucidation of the polymencharide from the steads of this important plant, Phaseolus sunge.

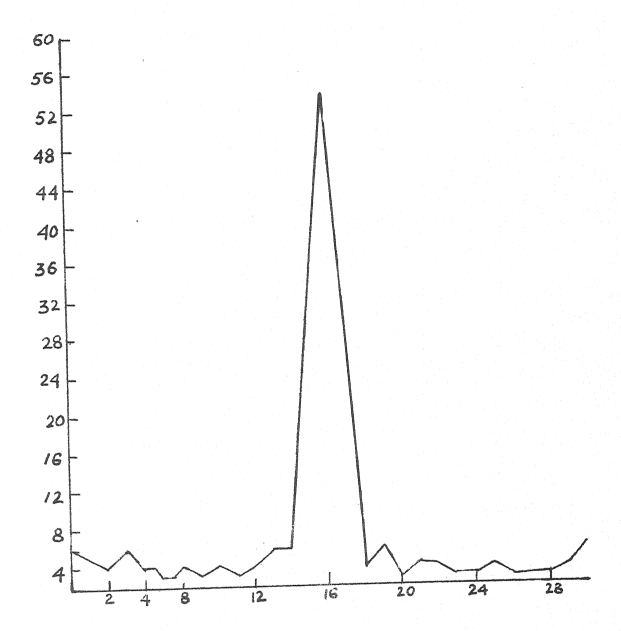
EMPLOYED STREET AND STREET OF MASSIONS BUTCO

111.2.1 RESULTS AND DESCRIPTION

A new weber soluble polymercharide has been isolated from the defetted seeds of P. number by extracting with 1% acetic acid and precipitating with emense of othered. The polymercharide was repeatedly purified till the seh content reduced to minimum. The homogeneity of the polymercharide was checked by a

- (1) Proctional Procipitation .
- (11) Zone electrophomesis , and
- (111) /cotylation despetylation.

into three fractions by fractional precipitation with different volumes of etherol. All the three samples were enelysed question to the polyacocharide to the calquad polyacocharide showing the polyacocharide to be homogeneous.



Segment number Fig. (1)

The portion of the polysaccharide was caparated by zoneelectrophorous method in borate buffer (pi 9.3). After completion of the experiment, a plot of the absorbance against segments
numbers should only a single sharp peak indicating the polysaccharide to be homogeneous. (Fig. 1)

The homogeneous polysoccharide was ecctylated with scatic anhydride and sodium scatato. The acetylated product showed optical rotation, $[\kappa]_0^{25} : 20.5^\circ$ (in chloroform, C, 0.88%). On descriptetion, it gave a polysoccharide having the same optical activity as the original one. Thus it confirmed the homogeneity of the polysoccharide.

111.2.2 The polysaccharide was alouly soluble in water. [2] 20 . 72.20 (in water, C, C.46), ash content 0.625. The polysaccharide was found to be free of nitrogen, sulphur, and halogens. The methoxyl, uronide and acetyl percentages were found to be negligible.

Likel. The complete acid hydrolymis of the polymorphic mily with analysis of the hydrolymate revealed by the passance of two sugars, D-galactome and D-mannose. The identity of the sugars was confirmed by their specific optical retations, preparation of their crystalline derivatives and co-chromotography with authorite samples.

The quantitative estimation of mono-sechanide components by periodate oxidation, taking ribose as a reference sugar, showed that galactose and menoose are present in the malar ratio, 1:4 in the polysaccharide.

The graded hydrolysis of the polysaccharide with 0.050 sulphumic acid and subsequent paper chromategraphic analysis of the hydrolysate, taking out at various intervale, revealed that galactose was liberated first followed by the liberation of mannose This shows that mennose units are linked tegether forming the backbone (main chain) of the polysaccharide and most of the galactose units are linked as terminal groups. The easy liberation of galactose units indicates that most probably they are linked to the main chain at peripheri through ~ linkeges.

method using dimethyl sulphate and alkali⁴⁴ followed by Furdie's method using dimethyl sulphate and alkali⁴⁴ followed by Furdie's method giving a methylated polyacocharide, [2] ²⁵ • 40.4° (in chloroform, C. 1.35), Cite, 46.4%. The complete hydrolysis of the methylated polyacocharide and paper chromatographic analysis of the hydrolysate in solvent (A), revealed the presence of four methylated sugars. The methylated sugars were separated on a pre-parative scale by chromatography on thetman No.3 filter paper. The following methylated sugars were identified.

- (I) 2,6-01-0-mothyl-Degalactores
- (II) 2,3-01-0-mothyl-0-monnogo;
- (III) 2,3,6-Tri-0-mothyl-0-mornoge;
- and (IV) 2,3,4,6-Tetro-O-mosthyl-D-galactoco.

in thyloted sugar, I, had $\Pi_{\rm DIG}$ in solvent (A), 0.46, [\swarrow] $_{0}^{25}$ \circ 80° (in mater, G, 0.66). It formed 2.6~di~0~cothyl~0~galactose entities on transmost with ethonolic entities, n.p. 123-22°, [\swarrow] $_{0}^{25}$ \circ 12° (in ethonol, C, 0.66). On emidetion with broadne mater it gave/ \circ leptons, [\swarrow] $_{0}^{25}$ \circ 22° (in mater, G, 1.25)

which on treatment with phenyl hydroxine formed 2,6-di-0-methyl galoctonic acid phenyl hydroxide, n.p. 130°. Thus the above observations confirmed that the methylated sugar, I, is 2,6-di-0-methyl-i-galoctops.

Methylated sugar, II, was obtained as a syrup, $n_{\rm DEG}$ in solvent (A) 0.56, $[\ll]_D^{26}$ = 16.80 (C, 1.80 in water). It formed 1.4.6-p-nitro benzoate with p-nitrobenzoyl chloride, n.p. 191-92 $[\ll]_D^{26}$ + 630 (in chloroform, C, 1.26), which shows that the methylated sugar, II, is 2.3-di-methyl-D-mannage.

Methyleted sugar, III, R_{BIG} in solvent (A), 0.63 . []
- 12.2° (in water, C, 1.6) formed 1.4-bis-p-nitrobenzente, a.p.
186-88° []
- 32° (in chloroform, C, 0.46). On exidation with
bromine water, it gave a lactone, which on treatment with phenyl
bydrazine formed 2.3,6-tri-0-cethyl-i-monnonic ecid phenyl
bydrazide, a.p. 129-30°. This indicates that the methylated sugar.
III is 2,3,6-tri-0-cethyl-0-centhyl-i-monnone.

Hethylated sugar, IV, R_{DAG} in solvent (A), 0.98 (C) 25 · 120° (in water, C, 0.65). On treatment with ethenolic eniline gave 2,3,4,-6-totre-0-mathyl-M-phanyl-D-galactosylandne, m.p. 188-90°. Therefore the identity of the mathylated sugar IV, is established as 2,3,4,6-totre-0-mathyl-M-palactose.

The quantitative estimation of methylated sugars by the method of Hirst and Jones 46 showed that the sugars I, II, III, and IV were present in the molocular metho, I : 4 : 20 : 5.

The studies indicate that galactose units in the polysaccheride occupy terminal positions as non-reducing and groups from which 2,3,4,6-tetra-0-mathyl-0-galactose IV., arises on hydrolysis of the methylated polysaccharide. A large portion of III, 2,3,6-tri-0-mathyl-0-mannose (20 moles) indicates that the back-bone of the polysaccharide consists of mannose units limbed through 1 ->4 linkeges. Isolation of 2,6-di-0-mathyl-0-galactose (1 mole) made on idea, that one male of galactose unit per repeating unit of the polysaccharide is linked at position 1, 3, and 4. Detection of 2,3-di-0-methyl-0-mannose (4 moles) shows that four mannose units in the main chain per repeating unit of the polysaccharide are linked at position +6 in addition to +1 and 4- positions.

Allah Determination of terminal groups by periodate exidation and subsequent titration of liberated formic acid, corresponds to 0.1038 males of formic acid per 100 g of the polysaccharide, is supposed to consist of 30 sugar moisties of which 3 units of galactose form terminal groups. Considering such a repeating unit, the terminal groups were found 16.80% as determined by periodate exidation studies, which identical to that revealed by methylation studies (16.79%).

Link. The partial acid hydrolysis of the polysocaharide followed by paper chromotographic separation on preparative scale afforded six oligosoccharides. The following oligosoccharides were detected:

- 1. Mannototroso, 0. P.D. Mannopyranosyl. (1 \Rightarrow 4).0. P.D. Mannopyranosyl. (1 \Rightarrow 4).0. P.D. Mannopyranosyl. (1 \Rightarrow 4). D. Mannopyranoso.
- 2. Mannotriose, $0 = \beta = 0$ -mannogyranesyl= $(1 \Rightarrow 4) = 0 = \beta = 0$ -mannopyranese.
- 3. Epimelibiose, 6-0-β-0-galactopyranosyl-0mannopyranose.
- 4. Mannobloss, 4-0-6-C-mennopyranesyl-C-mannopyranese.
- 5. 6^2 \ll galactopyl mannoblose, 0- \ll \approx galactopyranopyl- $(1 \rightarrow 6)$ \approx β \approx mannopyranopyl- $(1 \rightarrow 4)$ \approx D-mannopyranope.
- 6. Galactobiose, 3-0-<->—galactopyranosyl-0-galactopyranose.

Cligosaccharide, (1), m.p. 233-32 (2) - 260 (in water, C. 1.2%), was crystallized from aquason othered. It was found chromatographically sums in those solvents systems F. C and B (Page 84). The complete said hydrolysis followedly paper chromate-

graphic analysis revealed the presence of only menaces units in the eligomaccharide corresponds to a totresecularide. The hydrolysis with the ensure, equivalent and the negative retation indicated that the menaces units in the eligomaccharide are linked through β —linkeges. Fartial acid hydrolysis yielded menaces, menachiose, and menactrices which were identified by their co-chromatography with the authorite namples. The periodste emidation revealed the liberation of 2.12 moles of formic acid with the consumption of 6.20 moles of metapormic date per mole of the eligomaccharide. On the basis of these experimental evidences, the eligomaccharide has been identified as $O = \beta = O = menacopyrenesyl = (1 \rightarrow 4) = O = menacopyrenes$

C. 1.85) was chromatographically pure in solvents (C), (G) and (F). It was shown to be menchydrate of trisaccharide on the basis of its equivalent weight, 264.8. Acid hydrolysis of the eligeneochemide yielded only menages. The anomaric configuration of non-wedge ing manages units were found to be by enzymic hydrolysis and negative rotation was found within the range of that reported for mannetries. Partial acid hydrolysis yielded menages and menagicae which were identified by co-chromatography with the authentic samples. The identity was also confirmed by the pariodate oxidetion data which showed the liberation of 2.10 moles of formic acid with the consumption of 3.3 moles of metapariodate per mole of sugar. Hence the eligeneocharide was identified to be 0.7 mb-mannepyranesyl-(1.44).0.7 meannepyranesyl-0-mannepyranesy.

(Fig. * 3).

Fig-2

Fig-Z

Fig-4

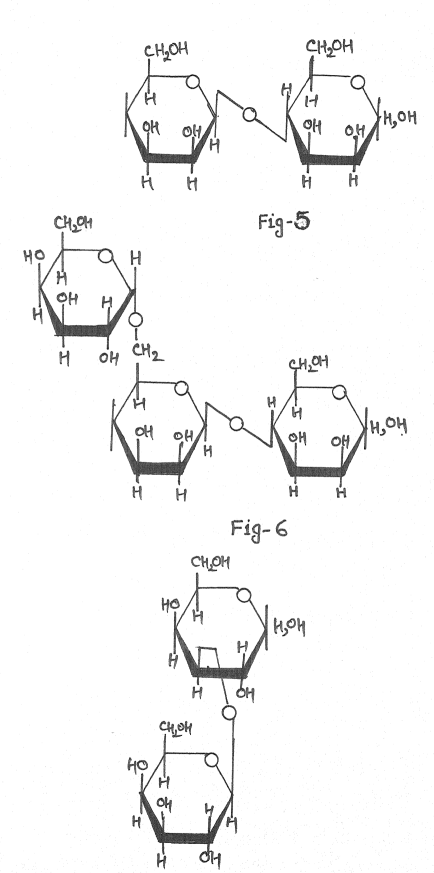


Fig-7

Oligosocharide (3), was isolated in crystalline form having the physical constants identical with those reported for \$-0-<-0-galactopyronosyl-0-connopyranose. It reduced Fehling*s colution and Tollen's respent having m.p. 200-010. 22. 120.40 (in water, C. O.48%) and was found to be a mingle entity by paper chromatography in three different solvent systems (A), (B) and (C). The paper chromatographic analysis of the completely hydrolysed sugar revealed the presence of galactoss and manness. The quantitotive estimation by the method of Hirst and Jones " showed the molar ratio to be 1:1 between the two sugars in the eligosecharide. The equivalent weight, 174.2, showed it to be a disappharide. The periodete emidetion studies efforded the liberation of 2.2 moles of formic sold and consumption of 5.24 moles of pariodate par mole of the diseacheride. The liberation of 3.2 moles of formic acid from the disappharide indicates that there is 1 -> 6 linkage between galactoss and manness units. As the disaccharide could not be hydrolygod with emulsin, it is inferred that calactose and mannes have & .linkege between them. On the basis of above evidences the oligosaccharide was identified to be epimeliblese, 6-Q-<-D-cologtopyranosyl-O-mannopyranose and identity was further confirmed by co-chromatography with an authentic mample (Fig. - 4).

of formic acid with the consumption of 4.22 makes of metaporiodate per make of the sugar. Hence the alignmentation was assigned the structure, $4-0+\beta=0$ -mannopyranosyl-0-mannopyranose. The identity was confirmed by co-chromatography with an authentic sample (Fig. + 3).

Oligosaccharide (5) was crystallised from ethanol, m.p. 266° - 268°, [] 32 + 90.0° (in water . 6, 0.46). It was shown to be a single entity by paper chromatography in solventerstans (G). (C) and (8) (page 84). It reduced Pehlings solution and Tollen's reagent The complete said hydrolysis of sugar and subsequent paper chrom. tographic examination revealed the presence of calectone and manager. The quantitative estimation by the method of Hirst and Jones should that galactose and mannose are present in the oligosaccharide in the ratio 1:2. The equivalent weight . 262.8. showed it to be a trisaccharide. The pariodate emidation studies showed the liberation of 3.18 males of famile acid with the consumption of 6.30 moles of metaperiodate. Partial acid hydrolysis fellowed by paper chromatographic examination showed the presence of mannebless and epimelibese besides galactose and mannose. Their identity was confirmed by co-chromatography with their authentic samples. The olicomecharide was, thus identified as 0. - alastopyranecyl- $(1 \rightarrow 6)$ -O- β -O-mannopyranosyl- $(1 \rightarrow 4)$ -O-mannopyranose. (Fig. - 6).

Cligosaccharide (6), $[\infty]_0^{30}$ + 152° (in veter, C, 1.26) was shown to be chromatographically gure in solvent system (G) (page 84) Acid hydrolysis showed the presence of only galactone units and its equivalent weight, 173.4, corresponded to a herose disaccharide. It could not be hydrolysed with exulpin. The periodote swidetion should the liberation of 1.66 males of female sold and the consumpt

tion of 3.12 moles of metaporiodate per mole of the sugar. The eligosecheride is, therefore identified to be 3.0- <-0-galactepyranosyl-0-galactess. (Fig. + 7.).

Link. On the besis of the regults obtained so for particularly from the methylation studies, graded and partial acid hydrolysis, the following valuable informations could be derived.

- (1) The main chain of the polysaccharide consists of β =(1 \Rightarrow 4) linked manage units.
- (11) One galectose unit per repeating unit of the polysecondride is 48650 also linked in the main chain through $\beta *(1 \Rightarrow 4)$ -linkage.
- (111) Galactose units form single unit branches linked to the main chain through <-linkages.
 - (iv) \ll =(1 \Rightarrow 6)=linkages between galactose and mennose units and \ll =(1 \Rightarrow 3)=linkage between two galactose units are present in the side chain only.

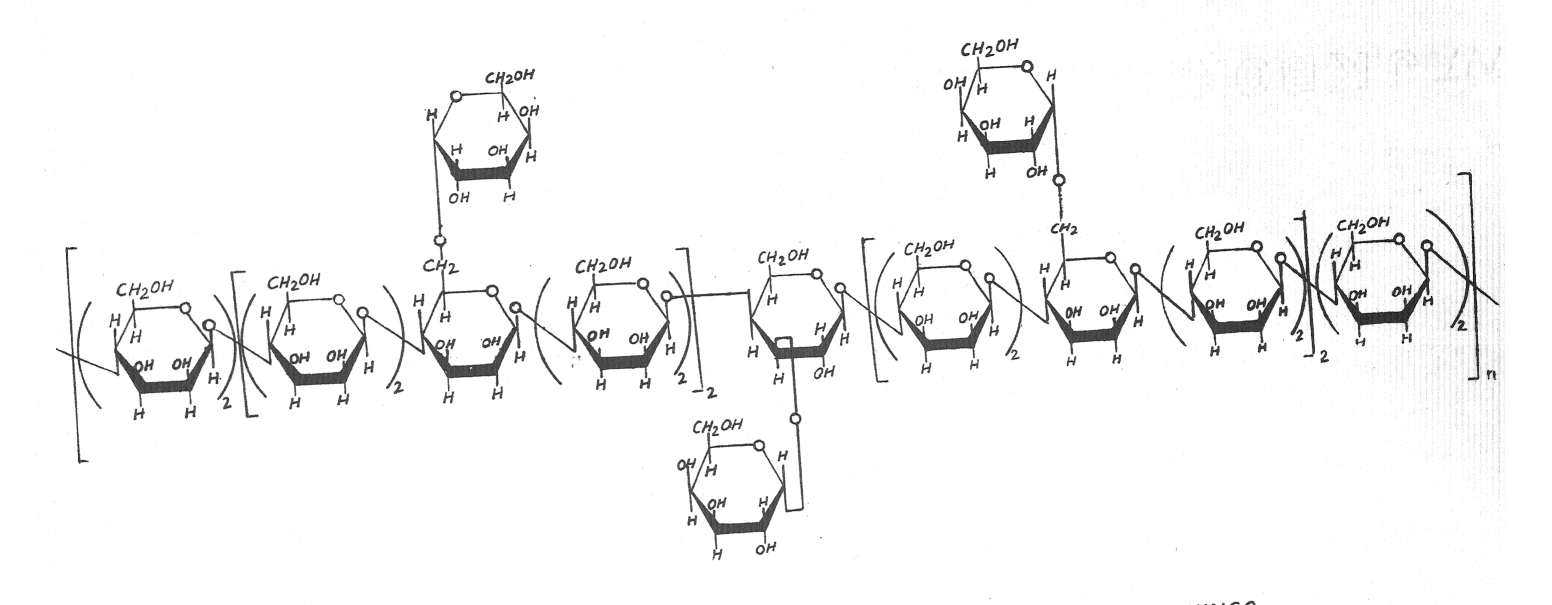
Taking all the experimental evidences into consideration together with the structures of different oligosoccharides, the following most probable structure has been assigned to the polyester-haride from the seeds of imagedus mange.

Galp = 3-galactopyranosa

The above structure contains 30 units of homes memosaccharides per repeating unit, which fully explains the formation of
oligosaccharides as obtained by partial acid hydrolysis and agrees
well with the analytical data of the polysaccharide. The detted
and doubly arrowed detted lines show the probable made of fission
of linkages during the partial acid hydrolysis. The arrowed
detted lines indicate secondary hydrolysis.

The polyentcharide such as described above about consume 34 males of metaparisdate with the liberation of 3 males of femile acid per reporting unit of 30 sugar units. The netwell consumption of periodate and liberation of formic acid have been determined to be 34.4 moles and 5.04 moles respectively per repeating units of polysaccharide which are in close agreement to the calculated values.

completely ruled out but they are less probable because the xxxxformation of eligosaccharide as obtained in the present case might
not be possible.



STRUCTURE OF POLYSACCHARIDE FROM THE SEEDS OF PHASEOLUS MUNGO

Paper chromatography was performed at room temperature by descending technique on whether No.1 filter paper unless stated otherwise using following solvent systems:

(A)	a-Butanol - ethanol - water	(51114)49,90
(B)	n-Butanol - acotid scid - water	(411.15) ⁹¹
(C)	a-Dutanol - iso-proponol - woter	(11:1:5)52
(D)	Bensone - ethanol - weter	(169147115)58
(8)	Butanono - woter	(10:1)54
(P)	Sthyl ogetote - pymidine - weter	(10:4:3)55
(G)	Sthyl acototo - pyridino - weter	(21112)96
(11)	n-Dutanol - othenol - water	(40:11:19)57
(1)	a-Dutanol - pyridino - water	(6:4:3)50

The spot were located by sproying the chromatgram with aniline hydrogen phtheists ond heating it at 110-20° for 10-15

minutes. Spectrometric determination were carried out by a modification of phonol-sulphuric acid method⁶⁰. Elett-Summerson phicologyric colorimeter was used for measuring the absorbance.

111.10 ISOLATION OF THE POLYBACCHARIDE

The dried and grushed seeds (2 kgs) were extracted successfully with petroleum ether (60-80°) and ethanol. The polysecharide extracted from the outracted seeds by repeating the process as given on page 31. A colourhess fibrous precipitate of the crude polysecharide was obtained. It was filtered, washed with absolute ethanol and dried in vaguum at room temperature (39 g. ash 2-135).

111.9 PERIFICATION

The dried grude polymersharide was dissolved in distilled water (2 litres) containing 15 exetic axid with constant stirring The solution was filtered and was added very slowly to ethanol (8 litres) with constant stirring and kept over-might. The precipitated polymersharide was filtered and the above process was repeated four times, to get a white fibrous mucilage, (26 g. ash 0.6).

III.11 HOMOGENEITY OF THE POLYSACSHARIDE

The homogeneity of the polysaccharide was checked by following methods.

III.11.1 (a) Promitional Promipitation

The pure mucilioge (4 g) was finetionally precipitated into

two fractions (Reaction 1 and Reaction II). Soth the fractions along with the original polyeaccharide were hydrolyped and quantitatively estimated by the usual way as described on page 32. The ratio of galactose and mennose in both the fractions was found almost the same (114) indispting the purified polyeaccheride to be homogeneous.

111.11.2 (b) Zone - Electrophoresis

Polysaccharide (300 mg) was taken for zone electrophoresis and similar procedure was adopted as described on page 34.

The corrected absorbence readings (Table - 1) so obtained were plotted against the distance from the anode, that is segment number which showed only one sharp peak indicating the polysacchantide to be homogeneous.

III.11.3 (c) Acetyletion and Descriptation

The pure polysaccharide (1.5 g) was mixed thoroughly with anhydrous sodium scetate (10 g) and the mixture was suspended in scetic anhydride (30 mls) and further process was repeated as on page. The scetylated polysaccharide (1.1 g) was obtained having $|\nabla ||_{D}^{25} + 28.5^{\circ}$ (in chloroform, C, 0.88%).

The dried acetylated polyenopharide (0.8 g) was dissolved in acetone (30 ml) and 50% potensium hydroxide solution (30 ml) was added to it. The acetylation was carried in the usual manner as given on page 33 * The descetylated polyenopharide (0.3 g) having [2] * 71.5 (in unter, 6, 0.61%)*

Jerans Ioe	Lost reading of clubs	Mana Jeth Sending		Absorbase
	25		3.0	0.000
2			2.0	0.004
3	24			0.006
•	24			0.004
5	24			0.004
				0.00
	22		1.0	0.002
			2.0	0.004
9	23		1.0	0.002
	24			0.004
11	23			0.002
12		22	2.0	0.004
13				0.006
14				0.000
			37.0	0.034
16	40		27.0	0.034
17				0.032
			2.0	0.003
	23		3.0	0.000
	23		1.0	
21	24		2.0	3.004
22				0.004
28			1.0	0.002
	23	22	1.0	0.002
	24		2.0	0.004
	23		1.0	
27			1.0	
	24		1.0	0.002
			2.0	0.00
30	2		4.0	Call

Absorbance was measured on 5 ml portion of coloured columns.

The original polysocthomide, $[\prec]_0^{25} + 72.2^5$ (in water, G. 0.65) and the polysocthomide obtained after description had almost the identical specific rotations indicating the homogeneity of the polysochamide.

111-12 ASS CONTEST

The dried polygoccharide (0.2~g) was ignited in a cilicatrucible previously heated to a constant weight. After ignition, the crucible was cooled in a deglerator and weighed. From the weight of residue (0.0014~g), the ash content was calculated to 0.62%.

TXI.13 MAYSIGAL AND CHEMICAL EXMEDIATION

It was a fibrous white powder, very light in weight, slowly soluble in water, $[\le]_0^{25} + 72.2^6$ (in water, C, 0.66). For the purpose of optical rotation, the solution was filtered through a sintered funcel to get a clear solution and the amount of polysecheride in the solution was determined colorimetrically. The polysecheride was found to be free of nitrogen, sulphur, and helogens. On treatment with fohling's solution, it formed an insoluble copper complex but did not reduce it.

THE 14 HAMENATURE OF THE SUCKES

The polysoccharide was expained for free sugare by applying three spots of its solution in water on a strip of thether Newl Silver paper (15 x 45 cms) and developed in solvent (A) as described on page 36. The three summy respect magnificance and and

twichlowoocotic acid⁶¹, aniline hydrogen phthaloto⁶⁷ and eilver sitrate in acotone followed by ethanolic acdium hydroxide⁶² on three different strips of above paper showed no spot, hence it showed that the polygocharide did not contain any free sugar-

111.15 METHONYL CHOUPS DETERMINATION

The percentage of methoxyl groups was determined by the method of Belcher, fildes and Mutten and was found to be megligible. (0.84%).

112.16 ACRIVI. CHAIPS DEFINATION

The method by Belchez and Godbert⁶⁴ was followed for the determination of acetyl group percentage with and without mucilage which was found insignificant (0.98%).

111.17 URONOR CONTENTS DEFERMANCE

The uronide contents were found to be negligible by the semi-micro method of Sakar, Foster, Siddiqui and Stacey 65.

III.18 INDROLVAIS OF ROLVEACELANIDE AND DETERMINATION OF

The purified mucilage (1.5 g) was dissolved in 2H sulphuric sold (100 ml) and was hydrolysed on a water-both for about 24 hours. The hydrolysete was neutralised with barium carbonate, filtered and concentrated under reduced pressure. The hydrolysete was examined

paper chromotographically for monosecherides.

111-18-1 (a) Paper Chromatography

The spots of hydrolysate were applied on two sheets of the trans No.1 filter paper. The papers were developed separately in solvents (A) and (B) by descending unidioansional technique. The chromatograms were six-dried and sprayed with uniline hydrogen phthelete. On heating them in an oven at 130° , each chromatogram showed two spots. The B_{g} and B_{G} values of the two spots correspondenced to D-galactoge and D-magnages as given in the following Table.

Sug-2 identified		ent (A)				
D=Galactosa	0.00	0.07		0.10	0.16	
D-Mannooo	0.12	0.11		0.20	0.2	
Shared Shirth an approximate about a story of colored a single fact the first star for the	Name of the Control o	and a second	eron parametra a combo visco de com		n sakerakinga king tilkalar bilakina habasi bila	MODELLA CONTRACTOR

G = 2,3,4,6-Tetro-O-methyl-O-galoctose.

The identity of the two supers was further confirmed by co-chromatography with authoratic samples of the sugers in the same solvent.

III.18.2 (b) Column Chrometography

A portion of hydrolyneto was discoved in a small amount of squaeous methonol (1:1) and adsorbed over a column of callulose (2 x 35 cms.). The column was left over-might and the separation was effected with solvent (A). Fractions amounting to 10 ml were collected and checked by paper chromatography with authentic samples of 3-galactose and 3-menness in solvent (B). The fractions 1-12 , containing same sugar were combined together and concentrated to give 3-menness. It was recrystallised from equeous methanol.

[4] 25 + 12.6 (in water, 5, 1.6 of per 100 ml of solution). The

(1) Delignnose phenyl bydrasone

following two derivatives were prepared.

Found

Glyon (Lit.)67

5.5. 192-940

139-200°

(11) -elennose p. n. glycosylamino bensele ecid

The derivotive was propaged according to the recent method of allia Ge.

Citon (IIIe.)

0-P- 180-010

1....0

The fractions 20-20 were mixed and concentrated to give 0-quiectose. It was recrystallised from equeous methanol, $\left|\mathcal{L}\right|_{0}^{20}$ o 79.2% in water, C. 0.5 g par 100 al of solution). The following derivatives were proposed :

(1) 0-Gelectose phenyl bydrasino

m.p. 133-540

Glvon (11t.)69

(11) N-p-Nittrophony: - D- Ghiortospionine

in a sacro-test tube, galactose (50 mg), p-nitrosmiline (50 mg), one drop of glocial scatic acid and four drops of methanol-water (0:1 m/v) were taken. The saluture was boiled for 8 minutes and kept over-night in a refrigorator. The crystalline product was filtered, weeked with cold ethanol, other and dried in vacuus. It melted at 218-19° after recrystallinetion from sethanol, 132.

111.18.3 (c) Thin - Loyer Chromatography

The plates were proposed from alusary of silica gel G in 0.18 solution of boxic seld and the spots of hydrolysets along with benzene : sestic seld : methanol (1 : 1 : 3) 11 and six-direct.

These plates were sprayed with smilling hydrogen phthalate respect on heating them at 120° in an electric oven two spots corresponding to segalactose and semanness were observed.

111.19 QUANTITATIVE ESTIMATION OF MONOS/CONVENDE

The polysoccheride (200 mg) was hydrolysed with 2% sulphuric acid (35 ml) for 24 hours on a boiling water-both and neutro-14sed with berium corbonate. Albose (20 mg) was added to the hydrolysate. The hydrolysate was applied on shatman No. 1 filter paper along with the guide spots. After developing in solvent (C), the strips corresponding to the suggest was quit with the help of guide acods and clutted. The simple was published with periodste and the

quantity of the monosoccharide estimated as described on page

						a e		
Calactosa	3.30	4.2	2.76				0.	
ilianno 00	13.60		1116	•	• 22	4.60		
Allogo	1.42	2.04		0		0.61	0.42	

^{*} Strongth of sodium hydroxide = N/124.8.

Assuming complete recovery of D-ribose the above results indicate that in the polysecheride D-galactose and D-manness are in the noise ratio of 1:4.

III.20 GRADED EMPROLYSIS OF THE KOLYSACCHARIDE

The polysoccharide (100 mg) was dissolved in 0.05% sulphumpic acid (20 ml). The hydrolysis carried out over a boiling water both. The hydrolysete, taken out at various intervals, were examined chromatographically, without removel of sulphumic acid using solvent (3) for the purpose of irrigation of the paper. Results are given in the Table * 4.

(in ploutes)		
10	•	
15	Gelectoce (Feint)	Too spots (very faint)
30	Gelectore	Three spots (Feint)
43	Galactoso	Three spots (clear)
60	Calactoso	No spots (clear)
120	Galactose + Hannese (wory faint	Two apots (class)
180	Galactoso + Mannoso	Some as above
240	Galactose + Mannose	Sand on expen

De-Galactose was found to liberate first followed by the liberation of Demanness. The easy release of Despherose leads to the conclusion that most of it is present as terminal group and not in the main chain of the polysoccharids.

III.21 AETHYLATION OF THE POLYSAKHARIDE

The polyecocharide (0 g) was methylated first by the method due to Ferikh, ingle and Shide 44 followed by Fundie's method 45 as usual described on page $43\ast$

The partly nothylated product was brownless have $(.30 \, .30)$ =0CH, $(.30 \, .30)$ = $(.30 \,$

1.3 g per 100 ml of solution) .

CATTOR OF ASSEMLATED SHAME

The hydrolymic of methylated polymechanide was carried by slight modification of method due to bouveng et-al⁷⁴. The methyla-ted polymechanide (100 mg) was discalved in 85% formic acid (20 ml) and rest of the process was carried out as described on page 4-4.

After separation on imateum No.1 filter paper in solvent (A), the chloroform shrometogram of syrup showed four spate after spray. Ing with antline hydrogen phthelete and drying at 1200. The Ross value of each methylated sugar was calculated in solvent (A) and was compared with that, given in literature as shown in the following Table =5.

Nothylated sugars Ligatified		99,50,91
2,6-01-0-cathyl-0-galactoss	0.46	0.44
2,3-04-0-coshyl=0-monnoss	0.30	0.04
2.3.6-7z1-0-cothyl-0-commoco	0.00	0.01
2.3.4.6-Tetra-O-mathyl-O-calactom	0.90	0.88

III.23 QUANTITATIVE ESTIMATEON OF METRYLATED SUCKES

114-23-4. The mostylated polysoscientide (200 mg) was bydrolysed as given about. In the bydrolysete glucose (40 mg) was added and The chromotograms were developed by the descending method using solvent (D) as described on page 45.

The sugare were estimated by alkaline hypotodite method as given on page 46. The results obtained are given in the Table - 6.

TABLE OF

	etion & Sugar						
	2,6-01-0- aothyl-0- galactosa	0.16	0.22	0.10	0.152	0.539	0.171
).	2,3gD1.eQe mathyleQe mannoso	0.64	0.88	0.72	0,608		0.684
170 180 180	2,3,6-Tri-0- mo thyl-0- mannoso		4.00	3.34	3.060	4.141	3.406
).	2,3,4,6-Tetra- 0-methyl-0- galactoss	0.43	1.	0.70	0.741	1.090	0.850
	Glucoco	1.25	1.76	1.042		1.004	1.270

The above results corresponded to an everyon maker ratio between A, B, C and D as 1 : 4 : 20 : 5. The methylated sugare were calculated as the methyl ethers of anhydroheness units i.e. $G_{10}^{11}18^{0}5$, $G_{2}^{11}16^{0}5$, and $G_{2}^{11}16^{0}5$ for tetres, tris, and discompathyl sugare respectively. An everage recovery of the methylated polymacohenide was found to be 99.90% assuming 100% resovery of glypotes.

TII-23-2 CHARACTERISATION OF BUTHYLATED SUCKES

The methylated polysoccharide was hydrolysed according to the method of Garege and Lindburg⁷⁶ as described on page 47*

The mixture of different mothylated sugars was recolved into five fractions on Whatman No.3 filter paper using solvent (D) Strips cotamining different individual methylated sugars were eluted with water. The eluctor were concentrated separately under reduced pressure and marked as fractions, I, II, III, E and IV.

III.23.3 Proction I

Solid . $R_{\rm BIG}$ in solvent (A), 0.46% . found : (He, 29.16% calculated for di-mothyl honors, CHe, 29.80%. It was pink becomes pot on approxing with p-mainidine hydrochloride, m.p. 116-18° $[<]_{D}^{25} + 30^{\circ}$ (in water, C, 0.6%). Lit., m.p. 119-20°, $[<]_{D}$. 46° (in water, C, 0.4%) \Longrightarrow + 34° (equilibrium value). It gaves 2.6-di-0-mothyl-0-galactose analide on treatment with etherolic malline, m.p. 120-22°, $[<]_{D}^{25} + 18^{\circ}$ (in etherol, C, 0.6%). Lit. To may be a solution of the solution

The solid (150 mg) was omidised with immains water and the product , after neutralisation with all wer carbonate, was distilled to give a symmp $\left[\left< \right|_{0}^{20} = 22^{\circ} \right]$ (in water, C, 1.26), iii. 79 for 2.6-di=0-mathyl= $\sqrt{-1}$ octors, $\left[\left< \right|_{0}^{12} = 49^{\circ} \right]$ (in water, equilibrium, C, 1.095). The lactons (50 mg) was allowed to react with phonyl hadrance (1 mole) in beiling ether for 15 minutes on removal of solvent and heating at 85° for two hours a drystelline product was obtained, m.p. 130°. Lit. for

2.6-di-O-mothyl gelectonic ecid phonyl hydraside, m.p. 1400.

111.23.4 Fraction 11

Syrup, $a_{\rm ENG}$ in solvent (A), 0.36, found : Gie, 29.445, calculated for dimethyl : Gie, 29.815, $[<]_{\rm D}^{26}$ = 16.8 (in water, C, 1.0%), Lit. 80 , di-O-methyl-D-manness, $[<]_{\rm D}$ = 16.0 (water).

The sugar (100 mg) was dissolved in pyridine. It was finally weahed with water and dissolved in chloroform. The insoluble portion was filtered out and the solvent from the filtrate was evaporated in a vacuum desicrator. The trude product was reservatablised from other, m.p. $191~92^{\circ}$, $[\checkmark]_{D}^{2d}$ + $63~8^{\circ}$ (in chloroform, C. 1.2%). Lit. The last openity of the chloroform of the colors of the colors of the chloroform).

111.23.5 Proction III

Symp. $R_{\rm DiG}$ in solvent (A), 0.83, Round : CHe, 41.1%, calculated for tri-cothyl hardes : CHe, 41.9%, $[<]_D^{25} = 12.2^9$ (in water, C, 1.6 g par 100 ml of solution), Lit. 82 , for 2.3,6-tri-0-methyl-D-mennose, $[<]_D = 10^9$ (in water).

The syrup (150 mg) was dissolved in day pyridine (6 ml) and treated with p-mitrobenzoyl chloride (500 mg) for 45 minutes at 60-70° and left over-might at room temperature. A saturated solution of codium bicerbonate was added drop-wice until no effer-venceone occurred. After adding water (15 ml), the product was extracted with chloroform. The extract was dried over sodium bul-photo, outside of solvent was taken and in vences and crystalliand

from petroleum ether, m.p. 186-68°, [] 26 + 32° (in chloreform, C. 0.4 g per 100 ml of colution), Lit. 83.84 for 1.4-bis-penitrobenzoete of 2.3.6-tri-0-methyl-0-mannose, m.p. 187-88° and [] + 33.0°. The syrup (180 mg) was exidised with bromine water and the product crystallised from acetone-petroleum ether, m.p. 81-82°, Lit. 53, for 2.3.6-tri-0-methyl-Y-(+)mannolectone, m.p. 82-83°. The lactone (75 mg) was boiled under refluxed with little amount of animal charcoal in ethanol and filtered. On cooling, a crystalline product was obtained which was recrystalline ed from ethanol, m.p. 129-30°, [] 25 = 18.6° (in water, C. 0.66), Lit. 86, for 2.3.6-tri-0-methyl-0-mannonic acid phenyl hydraside, m.p. 131°, [] = 20° (in water).

Ill.23.6 Fraction IV

Solid, $B_{\rm RiG}$ in solvent (A), 0.90, found : CHe, 51.4%, calculated for tetramethyl herose, CHe, 52.54%, $[\ll]_0^{28}$ + 120° (in water, C, 0.60); iit. 87 , 88 , for 2.3,4,6-tetra-0-methylese galactose, $[\ll]_0^{16}$ + 142° \Longrightarrow + 117° (equil.) in water (C, 1.1%), n.p. 70-72°. It gave red colour with aniline hydrogen phthelete. Its trootseat with algoholic aniline gave 2.3,4,6-tetra-0-methyle Hephenylesegalactosylamine, n.p. 183-90°.

111.24 PERIODATE OXIDATION OF THE POLYSACCHARIDE

III.24.1 (a) Liberation of Formic Acid 90 & Estimation of End Groups

The pelyeogenerics (5:0 mg) was discolved in water (50 ml) and in the solution, potentials chicalds (0.5 g) and 0.25% codius

metapariodate (49 ml) were added. The volume was made upto 140 ml with water. In a blank experiment potament chloride (0.5 g) and 0.25% modium metapariodate (60 ml) ware diluted to 140 ml with water. The oxidation was carried out in dark at room temperature as described on page 50. The aliquota (5 ml wave taken and ware titrated for liberated formic acid against 1/102.5 sodium hydroxide solution using methyl red as indicator. Results are given in Table = 7.

ted (72 hours) per 100 g of the polymencharide. The amount of formic acid liberated (72 hours) corresponds to 16,80% of anhydrous house units present as end groups. The titre values of alkeli at 48,60, and 72 hours indicated that one sole to formit acid was liberated per 1225 g, 1064 g, and 963,3g of the polymencharide respectively.

Time (in hours)	Volume of alkali used (in al)	(to eg)	ood lotel formic acid liberated
0	0.88	0.395	11.037
16	1.02	0.497	
24	1.16	0.520	14.574
36	1.32	0.502	
48	1.20	0.073	10.00
60	1.72	0.771	21.413
	1.90	0.832	23.07
72 96	1.90	0.032	23.675

III.24.2 (b) Consumption of Sodium Metaperiodote 85

The polysecoheride (250 mg) was dissolved in water (70 ml) to which 0.25% sodium metaperiodate (40 ml) was added and the total volume was made upto 120 ml with water. A blank was also prepared with 0.25% sodium metaperiodate (40 ml) diluted to 120 ml with water. The periodate exidation was carried out at room temperature as described on page 52. The liberated indicator 2 ml aliquots of minture and blank were titrated at various intervals against 0.0404% sodius thiosulphate solution using starch as indicator. The readings with the polysecoheride were substructed from the corresponding readings of controlled experiment to just the titre values. The results are given in Table * 8.

ASSESSED FOR

Mae (in house)	Hypo used (in ml)	Periodese consus and (in ag)	total paraodata consumed (in mg)
8 16 24 36 48	1.04 1.00 1.14 1.10	4.408 4.008 4.928 5.100 5.360	269.70 260.11 295.67 306.05 321.60
60 72 84 96	136 146 146	6.224 6.311 6.311	373.46 376.67 376.67

The amount of pariodate consumed (84 hours) commempede to the consumption of 0.7077 make of pariodate per 100 g of the polyanechemids. After 96 hours periodate amidisad solution (10 ml) was hydrolysed with 2% sulphumic seld (page 37). The hydrolyse to examined paper chromotographically for the presence of D-galactose and D-mannage but the chromatogram did not indicate the presence of any of the two sugars.

III.25 PARTIAL ACID HMDROLYSIS OF FOLYSACHIARIDE

The polymercharide (6 g) was suspended in water (500 ml) in a three necked flack, and stirmed mechanically and the name procedure was adopted as described on page 53 *

III.25.1 Exemination of the Precipitate

The precipitate was hydrolysed and identified similarly as described on page 54. The chromotograms showed three spots corresponding to Rg values of D-galactess, and D-manness which were confirmed by co-chromatography with their outhentic samples.

III.23.2 Exgmination of the Bydrolysate

Paper chromagraphic analysis of the bydrolysate over Whatman No.1 filter paper using solvents(A) and (B) and aniline bydrogen phthalate as a spraying reagent produced nine spots thereby indicating the presence of nine sugars.

III.23.3 Seporation of Oligosaccharides

The eyeap was dissolved in minimum quantity of water. It was separated by paper shromotography as described on page. The pagence was described from school and Six is inscribed of

oligosoccharides and two fractions of monosoccharides were obtained.

III.25.4 Examination of Praction I and Identification of Hannototracee

The fraction was pacrystallised from squarous ethenol map. $230 \text{--}32^{\circ}$ and $[\times]_{0}^{32} \text{--}27.6^{\circ}$ (in water, C, 1.2 g per 100 ml of solution). R_{lign} , 0.12, 0.02, and 0.09 in solvents (F), (C), and (3) respectively. It reduced Fehling's solution and smontagal silver nitrate.

The suger was hydrolysed with 20 sulphuric solid, meturelised with berium curbonate and filtered. The filtrate was
concentrated and examined by paper chromatography using solvents
(A) and (C). The chromatogram idmicated only one spot, commenpending to Rg value of memore. Thus sugar consists of only
mannose units. The equivalent weight of sugar was determined by
hypolodite method 46 and was found to be 337.5 which corresponded
to a tetrosaccheride.

The pariodate emidation of the eligosaucharide showed the consumption of 6.2 moles of the eligosaucharide matapariodate with the liberation of 2.12 moles of formic said per male of the eligosaucharide. The eligosaucharide was completely hydrolysed with emulain suggesting [3 eplycosidic limitages in the molecule.

All the show results indicate that the eligosectheride

is Bellongered Selimpelly selimpe the identification of the

sugar is well supported by its consecute found and reported in

literature shown in the following Table - 9.

	Constanta		20 portod	
1000	M•p•	230-320	232-34 ⁰ and 231.5-32 ⁰	(95, 91, 92)
	Optical rotation	[~] 32 - 27.80	[<] ₀ = 31° and = 30.7°	(55, 92)
	a _{llon} in solvent (F)	0.12	0.11	(55)
	a in solvent		0.19	(56)

III.25.5 Symplection of Praction II and Identification of Mannotriosa

 $R_{\rm den}$, 0.09 and $R_{\rm Glu}$, 0.35 in solvents(C) and (G) respectively, $R_{\rm Glu}$, 0.22 in solvent (F). The sugar was crystalised from ethanol, m.p. 164-66°, $[\sim]_{\rm D}^{32} = 18.8^{\circ}$ (in water, C, 1.83). It reduced Fehling's and Tellen's reagents.

The complete acid hydrolysis with 2% sulphuric acid, subsequent neutralisation with berium carbonate and examination by paper chromotography with an authoratic sample only one money secharide. Demands was obtained. The equivalent weight of the agger was found to be 264.8 by hypoiedite method 46. Partial acid hydrolysis of auger with 0.5% hydrochirole acid at 100° for 10 minutes regulted in formation of number and memblose which were identified by openhaneltography with their authoratic samples.

Periodate oxidation of the augar revealed that 2.10 moles of formic acid ware liberated and 5.3 makes of pariodate were consumed per male of the sugar. The sugar was completely hydrolyeed with emulsia suggesting that mennoes units are linked through β -glycosidic linkages. On the besis of above results the sugar was Adantified to be meanotriose, β witemposites β -Hanp-1 ---> 4Hanp which was further confirmed by its physical constants as shown in Table 10.

7.831.3 + 10

Constants	Found	Asported	Rofozónces
	264-66 ⁰	133 • 159 ⁰ and 214•15 ⁰ (anhydroue)	(91,93,95, 96,97)
Optical rotation	[×] 32 - 18.8°	[4] 0 - 15° 26°	(90)
a _{Clu} in Solvent (G) and Solvent	0.34 (F) 0.22	0.33 0.22	(55,56)

III.25.6 Examination of Praction III and Identification of E inelibiose

R_{iam} 0.16, 0.23, and 0.37 in solvents (A), (B) and (C) respectively. The sugar was recrystallised from ethanel, m.p. 200-01°, $[\propto]_0^{32} \circ 120.4°$ (in water , C, 0.46 g per 100 ml of solution).

Acid hydrolysis of the augus whith 24 sulphusis sold and noutrolinetion of the bydrolysate with benium combenets followed by paper chromatographic analysis with solvent (C) revealed the presence of galactose and manage in the sugar which was further confirmed by co-chromatography with authentic samples. The quantitative estimation by the method of Hirst and Jones 37 should the moler ratio to be like between the two sugars in the olige-seccharide.

The equivalent weight, as determined by hypoiodite method was found to be 174.2. The periodate emidation studies corresponded to the consumption of 5.24 males of metaperiodate and liberation of 3.2 males of fermic acid per male of the alignmentation. Thus there is 1 > 6 linkage between galactose and mannose units. As the alignmentate could not be bydrolysed with amulain, it was inferred that galactose and mannose have Mainkage between them.

ager or Jordvetivo	Constant	Round		Bologopeo
ipinelibiose	Bep.	200-010	203-03°	(92, 99)
esoidilemiqü	Optical rotation	(in voter)	[A] + 120.9 ^d	
			[∠]0 + 120.9' → 124.((10 m)	
üpimelibiese	a _{Clu} in colvent	(G) 0.60	0.39	(96)
Caasono	Da Do	1730	170-760	(100)

III.20.7 Exemination of Fraction IV and Identification of Example 100 and

 $R_{\rm Man}$ in solvents (A), (S), and (C) were found to be 0.27, 0.46 and 0.33 respectively. The sugar was recrystallised from methonol, m.p. 202^{0} , $[<]_{D}^{30} = 10.2^{0}$ (in water, C, 1.2 g. per 100 ml of solution).

routrolisation with borium carbonate and subsequent examination by paper chromatography showed the presence of manness units only. The equivalent weight was determined by hypotedite method 48 and was found to be 174.8.

The pariddete exidetion station chound the consumption of 4.22 moles of pariodete with the liberation of 2.14 moles of

formic acid per make of the sugar. The sugar was completely hydrolysed with couldn showing the presence of \$ *linkage between the mannose units which was also confirmed by the negative optical rotation of the sugar.

Thus the eligosecohemide is a disacchemide compose of D-mennose units linked through \$ -glycomidic linkegs. The sugar was identified to be memorises, 4-0-\$ -D-mennopyrenesyl-D-menno-pyrenesy. D-mennopyrenesyl-D-menno-tyreness, which was confirmed by proparing the essense derivetive, m.p. 2040 and co-chromotography with an authorite sample.

The constants of sugar are given in Table - 12.

Sugar or derivativo	Constant		laportod	0.5700000000
leanoblose	23+P+	202	202-049	(55,37,81, 91,92,94)
Mannoblose	Costaction [43		- 7° -> -9°	(35,61,91, 92,97)
//annoblose	R _{glu} io solvents(F) & (G)	0.52	0.52 0.65	(35,36)
lannobiosazono		204-059	200=06 ⁰	(33)

III.23.8 Examination of Progrice V and Identification of 6^2 - \ll -Galactomyl Hannoblose

R_{ion} 0.00 and 0.17 in polyants (C) and (B) respectively. The purposes and recognicalizated from 90% ethanol. The pupus partie tion chromotography revealed only one epot. $B_{\rm plu}$ in solvent (G) 0.33 ; m.p. 226-28° and $[C]_0^{32}+90-6$ (in water, G, 0.49 g per 100 ml of solution). It reduced febling's and Tollen's respents.

The complete eqid bydrolysic with 2% sulphuric scidents neutralisation with berium carbonets and shrometographic examination showed the presence of galacters and mennous in the sugar. The quantitative estimation by the method of Hirst and Jones 37 showed that galacters and mannose constitute the alignmentheride in the moler ratio of 142. The equivalent weight was found to be 262.8 by hypolodite method 46.

The periodate exidation studies reveiled that one male of of the eligosaccharide consumed 6.30 makes of mateperiodate and liberated 3.18 makes of formic acid. Fartial acid hydroly-sis revealed the presence of mannehices and epimalibiose besides galoctose and manness.

From the above observation, the sugar was identified to

be <-Galpal ---> 6-\$-danpal ----> 4-danpa. The observed data

were found in close agreement with the reported values in litera
ture as shown in Table - 13.

Constante		Reported	89/0301608
O.p.	27,4710	221-250	(72,92)
Optical KJ3	2 + 98+8°	[67] ³³³ ♦ 93.5°	(72,92)
		→ + 90.0°	
Acato in Solvent	0.33	•	(34)

111.25.9 Examination of Fraction VI and Identification of 3-0- ≪-Galactopyronosyl-D-Galactope

 a_{G_01} , 0.60 in solvent (G). The sugar was recrystallised from motherel. $[Z]_0^{30}$ + 15.20 (in water, C, 1.2%).

Acid hydrolysis with 21 sulphuric acid followed by neutralisation with berium carbonate and paper chromatographic examination showed the presence of galectose units only. The equivalent weight was found to be 173.4 by hypeiodite method⁴⁶ which corresponded to a disagehoride of hemose units.

Periodoto oxidation studies of the sugar revealed the communition of 3-12 moles of sodium metaperiodote liberating 1-08 moles of formic said per mole of the eligeneraberide.

It could not be hydrolysed with emulsin indicating the linkage between the galactose unit to be ---

The above observation identified the eligosacchemide to be $3\text{-}0\text{-}\infty\text{-}\text{sgalactopyranesyl-}\text{-}\text{sgalactope}$. The identity was further configned by preparing its essues, m.p. $235\text{-}37^{\circ}$, and scatte, m.p. $153\text{-}95^{\circ}$.

Sugar or dorivative	Constant	Round	Reported	Soferances
Galactoblose	Conteal rotation	[x] 0 + 15% [s	JD + 1200	(92)
Coallege	2.0			(92)
Apotato		10.00	157-35	(92)

111-25-10 Examination of Fraction VII and Identification of

 R_G , 0.82 in solvent (C), $R_{\rm ligs}$, 0.62 and 0.80 in solvents (A) and (B) respectively. The sugar crystellised from equeous methonol, $C_{\rm loc}^{32}$ + 60.2° (in water, C, 1.0%). It was identified to be D-galactose by co-chromatography with an authortic sample.

III.25-11 Exemination of Prestien VIII and Identification of

Delignous o

 R_g , 0.12 in solvent (A) and R_G , 1.08 in solvent (G), E_G^{22} + 12.6° (in water, C, 2.0). The sugar was identified to be D-mannose by co-chromatographic examination with an authentic sample.

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IV.1 In the present Chapter changed enguinetion of a starol and two flavonoids from the seeds of Dougua careto ling. has been described.

Carrot), belongs to the family unbellifered, is a hispid herb, 1 = 4 ft. high. Leaves 2-3 pinnes ; pinnes pinnetifid, segments nerrow.lengeoleto. Outer roys considered in fruit; bractacles many 3-fid and simple. Fruit 0-1 inch; bristles of secondary ridges long, glistoning white, consets at buse only, of the primary ridges and l., subglochidiste, corpo;hore undivided.

The common is entensively grown within the ages, as a cold weether crop. The plant is found wild in Europe, extends through west Asia, west-weeks to Kashair and along the Himsleyen Ranges within the temperate zone, and cultivated throughout India.

The seeds are useful in discoses of kidney and is dropey, hervine tenic. Sed given in to uterine pains. Antipolymourite substances from carrot may cure polynouritis in pigeons in those cases where the discose has developed quickly (within 20 days).

LVa2. The details of research work reported in the literature on this plant is given in tabular form on the next page.

lent P	0.340	Constituents	
1. Carrot		Antipolynouratic	(1918) ₃
2. Carrot (Red & white) Agietic	•	Vitenia A contant	(1934)4
3. Garrot	•	Amino ocid composition (deginino, Histidino, leoloucino, Joudino, Lysino, Mothionino, Shonylpaino, Throonino,	(1937) ⁵
5. Correct		Tryptophen and voline). Choline content	(1980)7
6. Gerrot	409	Pentothonic acid	(1952) ⁸ (1957) ⁹ (1965) ¹⁰
8. Carrot	Roots	Lycopersons Cerotein	(1939)
10. Cerrot	Roots	Manthophyll Maino ecid composition (Alemine, Aspertic acid,	(1948) ¹³
		Glutanic acid, /aparagina Cystine, Nistidine, leok Lucine, Methionine, Three Tryptophan, /rginine, Glu mine, Glycine, Serine an	ucino, onino, utam d Valine)
12. Garrot		Aldrin and Dieldrin Organic sold - Halic, Ci leogitude and trace of Succiaic and Pumaric sei	40•
J4. Comot	Laows		(1933) ¹⁶ (1956) ¹⁷

(Continued)

Lant	20000	Constituente	
15.G3570'S	Locus	Aldrin and Dieldrin	(1960)14
6.Carrot	Losvos	Crnithine Carbasoyl transferese inhibitor	(1963)18
7.602308		Witanin C	(1996)19
S.Corrot		Cyan1.d1ne	(\$962) ²⁰
9-Garros	Some	Disinfectants	$(1933)^{21}$
O.Garrot	Phloon	Cellulose	(1961) ²²
A.Corrot	Groon	Clycomidic bitter principle othereal oil, wax-like pets other soluble fot and proto-alkaledde - (Pyralidi and Coucles).	
22.Corrot	Julco	Total sugars, Protein, Fata, Capactate, Total acidity sch. Co. K. Na. P. Cl. Total Carotono β -Caroto Vitanin C.	(1963) ²³ Ine
23.Daucus carota		Ethereel oll,	(24, 25)
24. aucus carota		Vitamin V ₂ content	(1952)26
A. Paucus		β-Corotone content	(1952)27
26.0 apçus carots	12040	Aructoso, Clucoso, Sucroso	
27.0000 60208	Roots	Fiteliles lights and	(3971)29
28 •D ataque		Vatorio C	(3946)30

(Continued)

Plor		Verte	Gone & Lauren Vo	References
	Dougus Gazoko	Lonos	Lutedic=7-glucoside	(1950)31
31.	Daucus carota	Loavos	Hydrocarbons, Alcohols, and Phytosterols	(1980)318
\$2.	Daugus empota	Floors	Pignanta - Kaampferel-3 glucoside, Kaampferel-3- diglucoside.& apigania	(1963) ³²
	Daugus carota	Pruito	Sthemeel oil composition (Carotal, Gamanyl acotate and epomidibydromycaryophy	(1964) ³³
3.0	Daucus carote	Prušto	Volatile oil composition Proc acids, consisting of Isobutyric and Palmitic oc	(34) 5de
			ethers, terpense (pinene e 1-limolene	

leaves, flower, fruits, and seeds of the genus have been extensionally examined for various plant products, but no work has been reported on the study of polysecharide. On characal examinemation a polysecharide was isolated from this plant. But due to the paucity of the amount of polysecharide, the antire study has not been possible. Since no study on flavonoid compounds too has been reported from the seeds, therefore, author became interested to study thereoughly the chamical constituents from the seeds of Regume, Applicate.

TV. 3-EXTRACTION AND ESCLATION OF STEROL AND PLANSMOJDES

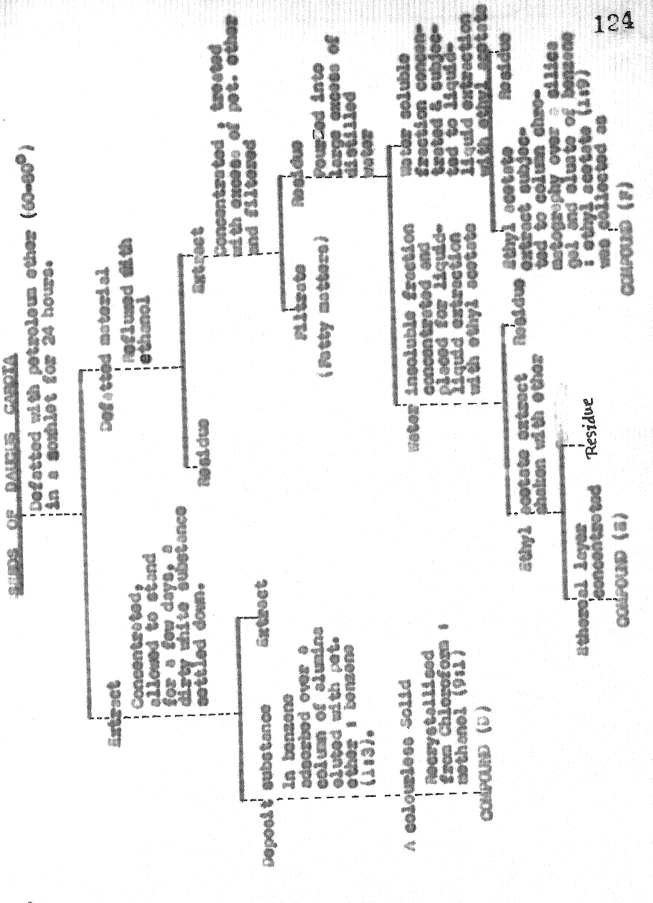
The seeds of <u>Daugus carots</u> were collected locally and identified for their authenticity in the Between Department of D. V. Postgraduate College, Crai.

ether (60 + 80°) in a southet extractor. This extract was concentrated and allowed to stand for a few days, when a dirty white substance settled down. The dayout was filtered from the extract dissolved in homeons and adsorbed over a column of neutral alumina. The column was eluted with the mixture of petroloum ether - benzame (1:3). The elucto was concentrated to give a substance which was recrystallised from chieroform - mothered (9:1) as a white flakes, compound (0), m.p. 138°. It gave characteristic Libermann-

The defetted meterial was extracted with ethanol (93%) on a steam-beth in several lots. The total extract was concentrated at reduced pressure to a brown viacous mass. It was refluxed with petrolous ether (40 - 60°) to remove the fatty metherial and resulting residue, still viacous mass, was powed into a large excess of distilled water with vigorous attraing. The water soluble and insoluble fractions were experated and successively subjected to liquid - liquid extraction, using petroleum ether, benzene, ethyl scotate and accome apparatoly.

The othyl nestete fraction of water insoluble part, was shown governi times with other in a separating funct. The ethoreal layer was esperated and the solvent was evaporated to drynoss where upon a light yellow substance was obtained. This on crystallisation from acctons * methonol (1:1) gave compound (2), m.p. 278° .

The acotone extrects of water soluble part was subjected to column chromatography over a silica gel G. The beasens - čthyl acotate (1:9) eluate of the column yielded a dark yellow coloured compound (F), m.p. 255-70°.



IDTAL

TV-S CHEMICAL STUDY OF GENERAL (D)

Compound (D), m.p. 130°, $[K]_{D}^{2D} = 36.6°$ (in chloroform), wes isolated from the seeds of Parkin satisfy as described on page [21* The compound was found to have molecular formula, $G_{20}^{11} g_{20}^{10}$, and soluble in petroleum other, benzane, chloroform, ethyl acetate, ethenol and methanol.

The compound gave all colour reactions specific for sterol, i.e., Liebermann-Gurchard reaction 38, Salkowski reaction 36, Tschugajaw reaction 37 and Kohlenberg's reaction 38. It also gave red colour with Noller's reagent 39. These reactions are specific for steroids and terpenoids. Since the compound did not produce any colour by Brieskorne test 40, showing the absence of triterpenoids. From the molecular formula and colour reaction, it is evident that the compound (D) is a sterol. It also gave positive test with tetra-nitromethams 41, indicating the presence of clefinic bend in the molecule, which is further supported by the posite at 845 and 805 cm⁻¹(Tri- substituted clefin) in its IR spectrum.

Con ago tyletics a memoratyl derivative, $C_{31}^{1}_{12}C_{2}$, maps 126°, $C_{31}^{2}_{12}C_{2}$ = 39.4° (in chloroform) and on benseyletics a memobensoyl derivative, $C_{36}^{1}_{34}C_{2}$, maps 142°, $C_{36}^{1}_{34}C_{2}$, maps 143°, $C_{36}^{1}_{34}C_{2}$

The NEW spectrum of the compound reveals the presence of five methyl groups contered at 0.48 (a) for methyl group at 0.43, 0.82 (4) for methyl group at 0.28, 0.92 (d) for two methyl groups at 0.25, 0.96 (d) for methyl group at 0.20 and 1.12 (5) for methyl group at 0.20.

On the bests of fewe-gains observations, the compound was identified to be β -mitosterol. The identity was confirmed by its mixed melting point, co-chromotography and the superimposition of its IN spectra over that an authentic sample of β -situatoral in the the compound (D) has been assigned the following structure.

β - Sitosterol (CINOUE) (D) .

- (111) Salkowski Rection ³⁶ . The chloroform colution of the compound on treatment with componitated sulphuric scid³⁶, gave a yellow colour which changed to deep mad.
- (iv) It discharged the colour of potassium permangements
- (v) tiollor's Reaction³⁰ . The compound gave a deep med colour with a few drops of thionyl chlorida (Prepared by adding 0.01% stannic chlordride in pure thionyl chloride).
- (vi) The compound was twested with concentrated hydrochloric end, along with a few drops of ferric chloride and the resulting minture eveporated to dryness. A red colour was produced when a few drops of water was added to the above mass.
- (vii) A white precipiote was obtained whom ethenolic solution of the compound was twested with ethenolic solution of digitoria.

TEMERAL ASSISTA

			Colculated for Coplant
G no	03.965		C = \$4.065
			H = 12.08%
	ular walght g method)	= 414	molecular weight = 414

TV-9 ACEDYLATION OF THE COMPOSED

So the compound (40 mg), fused sodium esstate (1 g) and esstic anhydride (5 ml) were edded and the whole reaction mistage was refluend for meanly 10 house over a gand-both at 140°. The

proction mixture was poured in ice cold water and precipitate, so obtained was weahed well with water, dried and recrystallised from chloroform - methanol (9:1) mixture, n.p. 125-27°, [S] 25 - 33.4° (in chloroform).

DETERMINATION OF ACTIVE PERCENTAGE

The percentage of ecetyl group in the scatylated derivative wes determined by the method of uleanbarger 44 as described by Delaher and Godbert 45,

	81.408		G	•	0157.5		
	11.465				11.40%		
Acoty	. 2020000000	11.025	Δο	o tyl			150

19.10 BENZOTLATION OF THE CUM-CUM

To the compound (20 mg), bensoyl chloride (2 ml) and four disps of pyridine were added in a Pyron stoppsred conical flack. The reaction minture was kept for 20 hours and heated over a water-both for six hours. The contents were cooled and poured in its pold water containing 25 aqueous sodium bicarbonate. The yallow realdes so obtained was washed wall with 25 sodium bicarbonate and folicated by water till it was free from the small of bensyel chloride. It was filtered, dried and recrystallised from chlorofers - ethenol (8:2) minture, m.p. 142°.

The escapeurd (30 mg) was discolved in hot absolute ethonol

(\$ mi) and treated with hot solution of digitaria (20 mg in 5 ml of absolute ethanol). The reaction minture was heated over a water-both for one hour, whereupon a white flocculent precipitate was obtained on cooling. It was weahed well with ethanol, dried and recrystallised from hot ethanol as a flocculent white solid, meps 215-47° (644, m.p. 210°).

IV-12 I-R. SPECIMEN OF COMPORED (D)

The following pools (on) in the IS exectsus (KBr) of the associated which observed by using Parkin-Siner Infra-good Spectro-

970, 945, 805, and 740.

IV.13 MAR SPECTEM OF THE COMPOUND (D)

The SEER spectre of the compound (D) was taken on Varian 4+40 Spectroseter, CDGL, so solvent and INS as reference.

			J	0.14	ŤÃ		
Outo	403						
0.02		(3	C	6		u)	
04.72	(4)			0.	40	¢,	ıą.
0.06							
1.12	46)						Selection of the select
LeO.	•2. _• 4) (;	'n.j				
	1.						

THE PROPERTY.

Hethyl group at C-13
Hethyl group at C-25
Hethyl group at C-25
Hethyl group at C-20
Hethyl group at C-10
Hethyl groups

TV.14 GESHICAL STUM OF COMPONE (8)

A light yellow coloured compound (8), m.p. 278° was included from the ethanolic extract of the seeds of <u>langua_gample</u> as described on page 122. The compound (8) having molecular formula. $G_{15}H_{10}\Phi_6$, was shown to be a single entity by paper chadmatography.

The otherolic solution of the compound gave following colour reactions.

- possine and bydrochloric ocid⁴⁶.
- (11) It gave an olive green colour with ethanolic fewric
- (\$11) It produced deep yellow colour with liquid emmonie and showed flowseconce under UV light 40.
- (iv) A yellowdeh trown colour was obtained on treatment with sedium hydroxide solution.
 - (w) It sould not be reduced with sedium berehydride 40.

the phone reservence suggest that the compound (8)

The sholding accounts only for C₁₅11₁₀C₂ which suggests that I have present as form

1001 the special series of this sholden is also supported to the sholden is also supported to the sholden is also supported to the compound of 257 as and 367 as with the compound series of the compound series and the compound series are postively.

The state of the solutions of those tydroxyl groups have been conigned to the basis of territors colour reactions, degradation and spectral station of the account. A free hydroxyl group at position -3

(1) A doup yollow colour was obtained when the methanolic colution of the appropriation was treated with electrics anythicalds.

which in the presence of citric onld gave a white procipitate 52-54

- (11) A deep yellow colour was obtained on addition of ethenolic boric acid and collium acetate reagents to the ethenolic polution of the compound ob.
- (111) A med colour was produced on treatment with zinc duct and hydrochlonic science.
- (Av) A bothochronic shift of 59 nm (> max from 367 nm to 426 mm) in the wisible region of spectrum of the eglycone was recorded on edulation of 1% ethenolic eluminium chloride to its ethenelic polution.

The compound gave plant colour with venillin hydrochloric sold reegent, shouling the presence of phloroglucinel unit in the ethucture, which compapends to the presence of free hydroxyl groups at position 45 and at 47 59.60. The presence of free bydrough at position =5 is confirmed by the following facts :

- (1) The compound produced an olive green colour with otherolle ferric chloride and a spot of this solution gave a yallan-wiolet fluomospense under V/ light⁶¹.
- (11) It give a yallow colour, on treatment with a solution of bonte seld and eltric seld in ocotons, which showed a yellow-Sah groom Sauomyauomea
- (411) The too depond spots of the compound on a filter paper produced bright fluctrosponce under UV light, when they were trouted separatoly with ethanolic solutions of aluminium chloride and zimponico osychiomido (1.44).

(iv) It gave red colour on treatment with Disgoth's #10000000

The presence of a free hydronyl group at position -7 is gonfigured by the fact that a bethachronic shift of 11 nm (λ_{max} from 267 nm to 278 nm) was observed on addition of fused sodium posture to the ethanolic colution of the compound52.

The mandadae fourth hydronyl eroup should occupy the position *4 in the psing B on the basis of fellowing facts :

- (1) A blue colour was produced on addition of sodium bilipprionute to the pink estution obtained by Shinode reduction
- (11) The mothyl other of the compound, on exidation with ndutral potacolum permangumate gave enisic acid . m.p. 1800 as one of the major enddetten products.
- (111) The wielble megion absorption maxima of the compound disappeared ($\lambda_{\rm mass}$ from 367 mm to 341 nm) when 0.002M sedium ethouside was added to its ethenolic solution, which showed a free hydroxyl at position +4" in conjugation with a free hydroxyl group at position 43 67,46,60.

The allowe enddenses indicated that the compound should hove the following structure:

This popposes the well known compound Managlerol.

This supposes by absorption tending of the solution obtained after Shinode reduction (\(\lambda_{\text{max}} \) 513 ms in sethemol. On its posits at 1665 cm \(\text{and 1615 cm } \) of the compound \(\text{of the instituted and 1615 cm } \) of the compound \(\text{of the instituted and the compound are confirmed by its mixed melting round and confirmed by its mixed melting and and co-chromotography with an authoratic sample.

TV-15 ISOIA TICK AND PURIFICATION

The compound (3), m.p. 270°, was isolated from the water insoluble frequion of ethanolic entract of the saude of Dansus.

twice hencemetry of the General

	l)			400		100	(41115 WV)	0.135

								di an anno constituti		arthu, arthu arthu
All the state of the		and the same of th			and the second second			E. 新山市 衛 JEEP 1	他在2011年2	0.52
			建设建筑建筑	30.				(60140)	TOTAL TOTAL	unites, applicate for the
No service of the service of	7 9 1.3	A TORSON STATES	THE PERSON NAMED IN	1987	distilluight uppressures				1990	

THE PARTY OF THE CENTRAL P.

	lound		Celculated for C15H10C6
		02,703	C = 62.93%
		3-42	H = 3.49%
Medicine		des union e 216	molecular weight = 206

172

The acetyl percentage in the acetylated derivative was determined by the method of Massenherger 44 as described on by Codeart and Delcher 45.

found

Colculated for CastaOa (COCHa)4

Acobyl group 36.90%

37.00

B B CONTROL OF THE CONTROL OF

the compound (2) (40 mg) was taken in dry scotons (20 ml) and anhydrous potnering carbonots (2.0 g) by refluciations or water-both for 22 locate. The momentum mixture was cooled, filtered and poured away careful for whomemore a vellocial mass was settled down. It was filtered, when well and recryptallised from ethenol, map.

PERSONAL DESCRIPTION DESCRIPTIONS

the methody's persontage in the methylated product wes determined by the puthed of Solcher, Fildes and Autten 72.

10.11.

Coloulated for GapHaOa (OCHa)

Notheryl group 31.2%

30.24

THE REPORT OF THE PROPERTY OF

OF THE CHICAL

The mothylated compound (20 mg) was exidised with neutral

potentium permanganate solution under reflux for four hours. The meaction mirture was cooled and the excess of mangamese dioxide was destroyed by adding sodium bisulphite to it. The resulting polution was acidified with dilute bydrochloric acid, whereupon a white compound separated. It was filteredend crystallised from ethonol, m.p. 1800. It was identified to be enisis egid by its mised melting point and co-charactography with an authentic sample is, 0.36 in n-butenol esturated with amonie ; spray - bromophenol blue solution).

20.19 UV and VISIBLE SPECIMA

WV and whalble spectre were recorded on Seckman model ou accompandamenter.

		begont	>						
	Uthanol.			267	*	367			
(44)				400	#	426	***		
(111)				243	*		11		
(24)	Echanol			400	(b)	341		25	
	l person			•		513	- 400		

Fellowing prominent pooks (co") were observed in the IR apoctrus of the compound.

3450, 3200, 1665, 1615, 1578, 1502, 1451, 1372, 1040, 625,

end 795.

1V.20 CHEMICAL STUY OF THE COMPOUND (F)

while entract of seeds of Q_{ABQDA} associate described a compound (F), s.p. $260\text{-}70^\circ$, having molecular formula, $G_{27}H_{30}G_{13}$. It was isolated from the seeds as described on page [22, and was shown to be single entity by paper chromatography.

The compound gave the following colour reactions :

- (1) It gave pink colour in Shinode reduction46, but did not give pink colour with hydrochloric acid only.
- (41) It gave on yellow openge colour with ethanolic
- (111) It produced yellow colour with liquid exmente and shewed yellow fluoroscence in UV light 48.
- $(\pm v)$ A yellowish colour was obtained on treatment with specime by drowide solution, which was stable on besting 73 .
- (w) with concentrated sulphuric acid, it gave intenses wellow colour with characteristic fluorescence 74,75
- (vi) to change in colour was observed on addition of

The above reactions suggest that the compound (f) is a flavore decimative peacecoing fallowing eheleten

It gave positive Holiogh's test indicating thereby the glycosidic nature of the compound. It is further supported by IR spectrum of the compound which exhibits the posits at 1120 cm⁻¹ and 1060 cm⁻¹ The exact nature of the glycoside was indicated by the identification and characterisation of the aglycone and sugar selecty absolute on each hydrolysis of the compound.

The pale yellow coloured aglycone, m_*p_* 349° has the meleschild formula, $C_{15}h_{10}c_{5}$ and responded to colour reactions (1 = 6) described obeton. The presence of this sheleton is also supported by the absorbation making of the compound at 269 nm and 336 nm. The eliminary three oxygen atoms may be present as three hydroxyl groups the compound formed triacetate and a trimethyl other on acetylation and mathylation respectively, confirming the presence of three hydroxyl groups. Thus the eglycone may be represented as below a hydroxyl groups.

The relative positions of these hydroxyl groups have been essigned on the basis of verious colour reactions, degradetion and spectral studies of the aglycome.

The eglycone on emidstion with newtral potageium permangenete gave a compound identified as p-hydroxy benzoic acid.

Aglycone of compound (F) Potasalum pasmanganete >

pathydroxy benzoic acid

This meetion shows that one hydroxyl group is present at position *4* of 3 ring of the compound. This was further confirmed by the following facts 4

- (1) when an excess of sodium bicarbonate was added to the solution resulting from Shinoda reduction of the compound, a blue colour 26,77 was obtained, showing the presence of free hydroxyl eroup at position =4.
- and a compound allowed a bathle compound a bathle compound allowed a bathle compound a bathle compound a bathle compound allowed a bathle compound allowed a bathle

(\$14) A bothophromic shift of 56 nm of Bend I (from 336 nm to 392 nm) without a decrease in relative intensity was observed by the addition of sodium ethomide to the methonolic solution of the compound. This shift is diagnostic \$7.60 for the presence of these bydromyl group at position *4.

(Ev) A single well defined posk (269 nm) of band li of the compound in ethanol also configured the presence of 4*-substintuent in the D-ring 70.

The position of 4 substituent in the $\frac{4\pi}{4\pi}$ is ring was limited supposed by this datas. The doublet 3-3.28 is the character for C-3 and C-3 protons while the other one at domestic data. As $\frac{4\pi}{4\pi}$ is for C-2 and C-5 protons, shielded by the C-4 and C-5 protons, shielded by the C-4

This degredation should the presence of free hydroxyl groups at spattions *5 and *7.

Phlorogluciani.

The processor of free hydroxyl group of position -5 was

(1) The eglycone (F) gave an arenge red colour with

Dimpoth's reagent (sertyl pyroborote) 65.

- (11) The aphytone gave bright yellow colour with methanolic sirconium oxychloride showing the presence of free hydroxyl group at position =3 ⁶⁰. The colour did not change on addition of strict acid showing the absonce of hydroxyl group at position =3 in the molecule ⁶⁰.
- twented with a solution of boric acid and citric acid in acetone, a yellow colour with a yellowish green fluorescence daveloped.

 This shows the presence of mothomyl or hydromyl group at postition as
- (iv) An ethanolic solution of the aglycone gave green colour with the ethanolic ferric chloride 47.
- (v) Sathochronic shifts of 46 mm in Sand I (from 336 mm to 382 mm) and of 9 mm in Sand II (from 269 mm to 276 mm) were elected by the addition of a few drops of ethanolic simulatum chicaids to the ethanolic solution of the aphycome. This showed a free hydroxyl group at position *5 of the aphycome 62,63.

The progence of free hydroxyl group at position =7 of the eglycone of the compound (F) was supported by the fellowing facts:

- (1) Pink colour was given by the aglycone with vanilia bydrochloride reagent, indicating the presence of 5,7-dibydroxy grouping 60 in the molecule.
- (11) A bothochronic shift of 9 ms of Sond II (from 20) ms

 (5) 173 ms | was observed on addition of a little funed podius accura

to the ethanolic solution of the eglycene, confirming the presence of free hydroxyl group at position *7. The eglycene also did not give any precipitate with neutral lead ecetate 61 showing the absence of ortho-dihydroxy grouping.

The supporting NEW data of 5.7-dibydromy grouping in the aglycene of compound (F) showed ad doublet at 3.95 and enother doublet at 3.55 which are indicative of proton at C-6 and C-8 in the ring A. It has been observed that flavones which contain 5.7-dibydromy grouping give rise to doublets (3 = 2.5 tps) in the range 3.5 to 4.05. A shorp singlet observed at 3.75 continue the proton N-3.

tiones, on the books of above observations, the aglycome of compound (F) has been assigned the following structure 4,5,7-talkydroxy flavone (/pigenin).

Paper chromatograph, of the sugar colution using mebutanel adotte octd - water (4:1:5 v/v) system revealed two spots with willies 0.16 and 0.20 respectively, suggesting the presence of a lactors and D-names. The Limitary of the sugars was confirmed by co-chromatography with me suthentic peoples of the sugars.

EV.23 PERTYTON OF CHICOGRAPH INCACE

The position of glycosidic linkego in the glycoside week determined by direct companison of the phydical and chemical properties with that of its splycose.

- (1) The glycoside did not respond to positive colour reaction with vanillin hydrophloric acid reagant whereas the aglycome indicating that the position "7 is involved in the plycosidic linkage.
- (11) The physocide did not give any shift in Sand II with funed condium exetate whomes aglycome of the compound gave a bethochmonic shift of 9 nm of Sand II tfrom 260 nm to 270 nm), confirming the presence of a free hydroxyl group at position -7 in the aphycome and absence of it in the physocide.
- (iii) The glycoside as well as the eplycome both gave positive colour test on addition of sedium bicarbonate to their respective Shinoda's reduction products. This indicates the absence of glycosidic linkage at position -4° in the glycoside.

Thus , it is only the position of in the aglycome at which both the sugars, Dogalactors and Domanness are attached. The pariodete original and also showed the consumption of 3.16 moles of pariodete with the liberation of 1.2 moles of fermic acid par mole of the physoside. It suggests that only one unit of each, galactors and manness is present in the molecular which corresponded to the molecular formula , $G_{22}H_{23}G_{13}$ of the physoside. The pariodete endotion studies also show that both the sugars are present in pyraness

form and are nutually limbed through 1 \Rightarrow 4 limbage in the discreteries. On partial acid hydrolysis of the glycoside by reflucing with 25 sulphuric acid an examined at different intervals by paper chromatography; galactose shound its appearance within can how indicating that galactose occupies the terminal position. After two and a helf hours of hydrolysis mannose appeared. The glycoside dissolved in hermal and hydrolysed with formic acid³⁴ for helf an hour. The aqueous hydrolysed with formic acid³⁴ for helf an hour. The aqueous hydrolysed gave a single spot by paper chromatography. The $R_{\rm g}$ value of this entity was not found to be identical with the $R_{\rm g}$ value of various noncencharides in diffement solvent systems $R_{\rm g}$ value of various noncencharides in diffement solvent systems $R_{\rm g}$ value of various noncencharides in diffement solvent systems $R_{\rm g}$ value of various noncencharides in diffement solvent systems $R_{\rm g}$ value of various noncencharides in diffement solvent systems $R_{\rm g}$ value of mannose.

The completely methyleted glycomide, on and hydrolysis, gave 2.3.4.6-tetro-0-methyle-0-galactoms and 2.3.6-tet-0-methyle-0-mannone which were identified by their R_{DIC}. It suggests that G, of mannose is involved in the glycomide formation with the approans. Finally, the glycomide was completely hydrolysed with smulain. This shows the processe of \$P-linkages between the giloc-tops and mannose and mannose and sunnose and splycome.

The above all evidences suggest that the compound (F) is Apigonin-7-0- β -D-galactopyranosyl-(1 \rightarrow 4)-O- β -D-mannepyranoside and may be represented as below :

TV DA TSULATION AND DURINGSATURE

The compound (F), m.p. 260-70° was isolated from the seeds of Daugus_Garata as described on page 12%.

W.28 POROGRAPHY OF YER CREENING

The homogeneity of the compound was checked by paper chromatography on thetman No.1 filter paper using following selvent systems:

- (1) a-Butanol acotic acid water (4:1:5,30:3:10 w/v).
- (11) Phonol saturated with water.
- (111) Acetic acid : hydrochlmeic acid : water (30:3:10 w/*).

In each case a single apot was objected.

HIGHENTAL ANALYSIS

		Colculated for Carling An
0 = 54.65		G = 54.5 46
11 m 5.10		11 = 5.00%.
	solcht = 594	Molecular weight = 594.

The glycoside (400 mg) was hydrolysed with %% ethenolic sulphusic sold (50 ml) on a water-both for 10 hours. The hydrolyset was cooled, solvent distilled off, diluted with water and filtered. The precipiets was dried in vacuum, crystallised from othyl costate - patrolous other (7+3) minture and finally recrystallised from tallined from matchinol to yield a pale yellow coloured compound

(aglycone), m.p. 349°. The filtrate obtained after removal of the aglycone was neutralised with barium carbonate, filtered and concentrated under reduced pressure to a syrupy mass.

IV.26 EXAMENATION OF ASSYCOME

It was soluble in ethenol, methenol, ecetone, pyridine and insoluble in petroleum other, beamens and water. It gave all positive tests, characteristics of flavonoids, as described on page—for the study of eglycone.

TV.27 CHROMATOGRAPHY OF AGLYCONE

The purity of the eglycone was checked on Whatman No.1 filter paper when a single spot was observed in each case using following solvent systems :

- (1) n-Butanol acetic sold water (4:1:5 w/v) 0.88.
- (11) Shenol seturated with water 0.95.
- (111) m-Greeol acetic acid water (50:2:48/w/v) 0.87.

ELEMENTAL ANALYSIS OF THE ACLYCING

				lated	
ı	66.72		6 0	(CaC)	
ž.					

AV.28 ACETYLATION OF ACENCENE

The eglycome (40 mg) was acetylated with acetic unhydride (50 ml) and pyridine (3.0 ml). The reaction minture was left

DETERMINATION OF ACTIVE PERCENTAGE

The acetyl percentage in the acetylated derivative was determined by the method of the emberger 44 as described by Godbert and Bolcher 45 .

Calculated for CashyOn(COCHala

Acotyl group = 32.69%

m 32.57/.

TO A STREET OF A STREET

The aphycome (40 mg) was taken in dry scattone (20 ml) and was mothylated with dimethyl sulphate (5 ml) and denhydrous potestion carbonate (1.0 g) by refluxing it on a water-both for 20 hours. The reaction mixture was cooled, filtered and powed over graphed ice whoreupon a yellowish mass settled down. It was filtered, weshed well and recrystallised from otherel, m.p.

DETERMINATION OF METHOMIL PERCENTAGE

The methodyl percentage in the methylated aglycone was determined by the method of Delcher, Fildes and Nutten 72 .

Pound

Calculated for Gasty Cocka) a

: 30.05

ss Linux .

TV-30 PCEARRIER PRINCIPATE OCTOATION OF THE METERS. STREET

The methylated algycome (25 mg) was emidised with newtral petassium permanganate solution under reflux for four house. The reaction mixture was cooled and the emeass of mangamess dismide was destroyed by adding sodium bisulphite to it. The resulting solution was acidified with dilute Bydrochloric acid, wherever a white compound separated. It was filtered and sacrystallised from ethonol, m.p. 170°. It was identified to be emisic acid by its missed selting point and co-chromatography with an authorite sample. The 0.36 in n-butanol saturated with samonias spray bromophenol blue splution).

31. DENTE CANADA DE STANDA

The ayrup obtained after the bydrolysis of the glycoside was demanded paper chromatographically using a-butanol - scatic sold - water (4:1:5 v/v) as immigating solvent system. The days looped chromatogram was sir-dried , aprayed with amilian bydrogen phthelate and an hasting at 120° for 10 minutes , two spots , by walues 0.16 and 0.20 were observed, which comresponded to lemphicators and D-mannoss propositively.

The Lieutity of sugars was confirmed by co-discussive states of the sugars was confirmed by co-discussive states and co-discussive states and co-discussive states and co-discussive states and

C. The second

The methyloted plycopide (30 mg) was hydrolysed with 34 methanolic sulphuric acid (30 ml) on a water-both for 4 hours under methanolic sulphuric acid (30 ml) on a water-both for 4 hours under methanol, the processor, and poured in distilled water. The processive was filtered, weehed well and recrystallised from methanol. The filtereds was nowhead with bories carbonate and consentrated under mader methanol was filtereds was neutralised with bories carbonate and consentrated under mader methanol processes to a light yellow coloured symme.

CASCA THE TRANSPORT OF THE PROPERTY OF THE PRO

the symup of the methylated sugars obtained as above was charactersproposed on whether No.1 filter paper using n-butanel - ethonol - water (5:1:4 y/v) as irrigating solvent system. The developed characterspropose was alr-dried, Sprysed with aniline hydrogen phthalate and heated to 1:20° for 10 minutes, whereupon two spots were obtained. The Arac values (Tag = 2,3,4,6-tetra-0-methyla-0-phinates) of the spots were found to be 0.80 and 0.90 which corner-pended to 2,3,6-tri-0-methyla-0-philatese manness and 2,3,4,6-tetra-0-methyla-0-philatese manness and 2,3,4,6-tetra-0-methyla-0-m

IV-33 PARTIAL PROPOLYSIS OF THE GINCOSIDE

Dulphanic science (2) no hydrolysed by reflucting although the particular of the property of the particular of the parti

The Sales Made

to do to cod.

IV.34 INCROLYSIS OF THE GLACOSIDE MIDI PORMIC ACID⁸⁴

The phycoside (20 mg) was dissolved in boiling cycloberanci (10 ml) and hydrolysed with formic acid (7%, 6 ml) by refluxing on a weber-both for helf an hour. The equeous hydrolysets gave a single apply by paper chromotography. The R_g value of this entity and not found to be disabled with the R_g value of various mono-calculations as different solvent systems. Further hydrolysis

The state of the s

(20 mg) was disposived in a minimum of themselves (25 mi) and (25

molecular weight of the compound	(1)	
for 15 ml eliquote of the reaction	60	0.58 al
mintum 0.010 modium hydronido s		
		3.2 81
For duck made of the chycoside		
makes of formic odd liberated	13	1.02
Holes of perfectate exercises		3.34

1V.30

The glycoside (20 mg) was dissolved in equeous ethanel end emulsia solution (25 ml), propored from elmonds 06 was added to it and the solution was kept at room temperature for four days. The hydrolysate after outrection with othyl ecotete. was concentrated to a syrup. The paper chronatography of the myrup in n+butenel - ecetic sold - weter (4:1:5 w/v) revealed the promote of two sports . R. O.16 and O.20, corresponding to gelectose and macross respectively.

WISIBLE SPECTRA OF THE CUSPORNO (P)

UV and whatble spectre were recorded on Deckman Model DU Spectrophotometer.

		and magant	λ_{mon}			SMft.
(1)	٨	+ 200mol		•		
(11)	A	+ Digitariol + NaDric	260	•	3.4	•
	٨	* Sthonol * AlCl ₃		•		10 . 45
	٨	+ Sthenol + NeOit				• 32
(4.)			200		3.35	•
(11)		e 200000 + 140/6	270	0		9 . 40
(ana)	D	* 44.44 ₃				9 . 46
(4)						8,96

14.38 IR SPECTRUM OF CONFOUND (F)

Pollowing prominent peaks (on") were observed in the IR spectrum of the aglycene :

3442, 3399, 1660, 1625, 1590, 1580, 1395, 1205, 1120, 1060, 842, and 710.

AV. 39 RESERVE OF ACENCINE OF COMPOUND (F)

1948 Spectre was recorded on Varian A-60 instrument using

	ઇ જ					
2.1 - 2	Las (a)	2000	at	C-2*		
	-28 (a)		at	(m)*	and	(See ja)
3.58	(a)	Protons	at	()=()		
* 3 8	(0)	Proton	36	C=3		
3.08		Paoton	9\$ (*		

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GIAMIN ... V

Mono

THE PARTY SHEET

Mal. The present Chapter deals with the Leolation and identification of water colubic monococcharides from the flowers of Linua unitationisms, Linua, commonly known as Alai, Tiel, Birl (Flam or Linused) belonging to the family Lineages.

Linua waitoticalmum is a annual horb, stam, 2-4 ft., orect, usually corymbosely branched above. Leaves linear or lanceolate, without stipular glands, sub-3-marved. Flower is broad cymes, blue or sometimes white, 1 inch across. Sepels evete, acuminate, 3-marved eglandular, margins white, ciliate or not. Stigmas linear - clavate. Capcule herdly exceeding the sepals a edges of valves ciliate.

The plant is cultivated throughout India upto 6,000 ft especially in the Denarce Division, in Bundalkhandand Sub-Mimalayan tract.

The plant is of high medicinal value Dried ripe seeds used as demule and in form of poulties, as poulties useful for gouty and rheum, smalling, used internally for irritation of genite wurinary system. Flowers used in hervine and condict tends. Oils mixed with line water used as application to burns.

From the survey of literature of Plant, the research work has been done till now shown in tabular fown on the next page of this chapter.

Lani	. epocioo	Porto	Gonati tuente	lo2opone0s
			Spinoble fibors	(1930)3
2.			Tombile fibers	(1933)
	raes:		Degradation and molocular size of flax postin	(1940)3
4.	Man	Plonto	Liberation of Plant fibers	(1936)6
6.	Flor	Zanta	Chamical composition plant fibers	(1969)
	Plan.	Elboro	Viadeoco, ortificial cotton and ortificial wool	(2941)8
7.			Flevonoids, querestin Keenpferol, epigenia genistein glycosides.	end
0.	SLOR.	Stoos	Separation of flam	(1936)10
9.	12.03	\$ 100 8	Postin A and Postin &	
10.			Poetin A and Poetin	(1941)13
		300dg	Structure of aldelica acid from flow sould murilogs	Me (1939)**
		9000	Bucklege, and fraction of sucklege	
		Soodo	14gnlas	(1961)16
		Soods	Bydrolytic studios o mucilago, Galacto wa	(1961) ¹⁷

(Concinued)

	3 (2003/96)	Vorto	Constituents Net	
	9101	Soods	Proportion of mucilage, characteristics of some polymomehamide	
10.			Four polyseccharides from lineard suchlage	(1970) ¹⁰
17.			Linonopin	(1952) ¹⁹
10.			Pet content	(1963)20
19.	14.58		throo fotby onldo	(1967)21
	F1.01		Hoophetide content	(1969) ²²
21.		Cotyledone	Sin glycoflowene Gwglycogide	(1969) ²⁴

grantable work has been done on the sood musilage and polymetable gide, and various other parts of the plant have been also investigated. But no work has been done on the flower of the plant.

Due to its medicinal importance it was consider worthwhile to grady the flowers of the plant.

The flowers of <u>Linux</u> unitationinum were collected locally and botomically identified in the Sotomy Department of D. V. Footgraduate College, Grai.

The crushed flowers were defetted with potentions other (60 = 60°) and extracted with cold distilled water. The whole lot was concentrated to a syrup which was examined for management rides by paper chromatography and column chromatography.

The two cheets of paper were propored and developed in Solutions (A) and (B). On appropring with amilian hydrogen phthelets each of the chromatogram showed the presence of five spots. The R_g and R_g values of these sugars corresponded to D-galactost. D-glucost, L-archinese, D-myless and L-rhomese respectively. These sugars were further confirmed by do-ahromatography with their authentic samples and on colluless column, the chromato-gram of above syrup was made which was run with the solvent (A) into different fractions. By: checking each elute on paper chromatography, total five fractions I, II, IXI, IV, and V were separated. The identity of each fractions was technic by paper chromatography on Westman No.1 filter paper with their suthentic samples. The further identity of each was confirmed by their m.p., m.mp., specific rotation and by preparing their derivatives.

Which Proction I, was expetallised from equature with 93-94°, $[<]_0^{37} + 8.6°$ (in water, C. 1%) gave pinkish colour with premiodine phosphote. It was identified to be i-shanness. Further identity was confirmed by proparing its premiodine phosphote declivative.

rection II, was erystallised from absolute ethanol, the 143-444 . [4] 0 + 17-0 (in ethanol water, C, 1-145). This fraction was identified as D-mylose, and further confirmed by proposion myloseson ducknotive.

Fraction III also erystallised from aqueous othenol, m.p. 150°. [] 39 + 101° (in unter, C. 15). From the above observations, the sugar was identified to be i-erabinose. Ito identity was confirmed proporing Learnbinose phonyl hydrorems doržvetivo.

Proption IV, orystallised from aqueous methanol, mep. 146° . $[\checkmark]_{0}^{21}$. 53°. On the books of the above results the sugar was identified to be D-glucoso.

Proction V, crystalliend from methanol, m.p. 165-46°, $\lceil \prec \rceil$ 29 + 80.8° (in woter, C, 15). All the above results indicated that the sugar was Degalactors. The identity was further confirmed by propaging its N-p-nitrophanyl-D-galactosylamine

V.3 EXTRACTION OF SUGARS

Actual defects with policious other (60 + 10°). The defector of the state of the st

The fellowing solvente systems were used for chromatography :

- (A) medutanol ethanol water (4:1:5)27:28.
- (3) m-Butanol acotic acid water (4:1:3)27.
- (G) Ethyl ecetate seatic sold weter (3:1:3)29.

Sprey RossonC

by satisfied states (0.93 g) and sathable acid (1.66 g) to water satisfied a letteral (1.00 ml). Aldohomomom give brown colour and altopositions size bright and colour with this reasons.

V-3-2 TOWNSHIP OF PAPER CONCOLTOGRAPHY

The goods of eyrup ware applied on the shoots of thatmen

No.1 filter paper. The paper were developed emparately in solvents (A) and (B) or by descending unidemphismal technique. The chromotograms were air dried and sprayed with amiliae hydrogen phthalote. On heating them in an even at 120°, each chromotogram showed five spote. The Sg and Sg values of the five spote corresponded to Degalactors, Deglucosa, Learnbinosa, Desyloss and Leghannosa as given in the following Table * 2.

Sugar Adontified	io dound	1000 440 1000 27,23	A Control	
	0.00	0.07	0.43	0.16
	0.00	0.00	0.18	
	0.11	0.42	0.20	0.23
	0.16	0.19	0.30	
4-21-22-00-0	0.5	0.30	0.00	0.37

G - 2,3,4,6-Tetro-O-mathyl-O-glucose.

the identity of the five augure was further confirmed by co-charactership with outbentic scople of the sugars in the same

AND THE STATE OF STREET

allege amount of above syrup was dispolved in a small amount of squares methanol (1:1) and admorped over a column of collider (2:s 35 cms). The column was left over-might and the apparention was effected with Solvent (A). Proctions amounting

to 25 al were collected and checked by paper chromatography with the authorate employ of different sugars.

Proctions I = VI containing sums sugar ware combined together and companies to give i-channess. It was expetallised from equeous etheral and gove expetals of *i-channess hydrote. Its m.p. and m.m.p. with an authorite specimen was found to be $93-94^\circ$, $[<]_0^{37}$ + 9.6° (in water, C. 16), Lit. 31,33,34 .

It gave pinkish colour with p-anisidine phospheto.

printedian Phosphoto - It was proposed by dissolving printedian (0.1 g) in phosphosic acid (4 ml. sp. gr. 1.75) and diluting the solution with ethanol (100 ml). The procipitated production phosphoto was filtered out, dissolving in minimum amount of water and mind with ethanol (100 ml). This solution was acidified with phosphombs acid (2 ml; sp. gr. 1.75) and mined with the first polution.

La Desylogo

Frection 8-14 were mixed and concentrated to give 5-sylese.

It was recrystallised with obsolute ethanol, s.p. 142-44°, not
depresent by mixing with an authoratic sample of 5-sylese.

[-] ** 17-5° (in water, G. 1-146).

Mylocomono - In a tout tube mylose (190 mg), phosyl bydro-

sine hydrochloride (300 mg) and sodium eastere (200 mg), discolved in water (7 ml) and heated on a balling water-bath for 30 minutes. Precipitate of the operane started appearing after 7 minutes. The flocquient precipitate was separated with water and recryetablished from 30% ethonol, m.p. 160-61° equal to the the authoratic started sample.

III. LeAponinom

Praction 15-21 containing sum augar, were combined together and concentrated to give i-exchinese. It was crystallised from equature othered, m.p. and m.m.p. with an authorite comple who 150°. [] 29 + 101° (in water, C, 1%), 140, 31,32,34

Le/mobines thenyl Sydrozobe . The sugar (200 mg), phonyl bydrozine bydrochhomide (400 mg), crystelline sodium acetate (600 mg) and water (6 ml) umme heated in a loosely stoppered test tube on a water-bath for 30 minutes. The tube was them cooled to reem temperature, the crystelline ceasons filtered out, weeked with distilled water and recrystellined from aqueous ethanol, maps 163-64°, lit. 36,37.

AV. D=60.000000.

Praction 23-27 were aimed and concentrated to give D-glucose. It was recrystallised from equators methanol, m.p. 146°, $[<]_0^{21}$. It was recrystallised from equators methanol, m.p. 146°, $[<]_0^{21}$

Phonyle-Gugogasono - Sugar (40 mg), sodium apetate (40 mg), 3 drope of phonyl hydranine, 1 ml of water and 3 drope of glocial costic cold were missed. After boiling the whole contents for 30 minutes, 2 ml of water was added and coaled, washed with water. The grystale of phonyle-Geglucosasses were obtained and recrystalized from dilute ethanol into medie form. This darke value has , m.p. 205°.

Proction 30-38 containing same sugar wars combined together and concentrated to give D-galectors. It was recrystallised from equators methods. Its map. $165-66^{\circ}$, $\left| \left\langle \cdot \right| \right|_{0}^{29} + 80 \cdot 6^{\circ}$, (in water, C. 1%) iii. 31,32 , map. $166-66^{\circ}$, $\left| \left\langle \cdot \right| \right|_{0}^{20} + 80 \cdot 2^{\circ}$ (in water, C. 1%).

were taken galactors (50 mg), p-nitrouniline (50 mg), 1 drop of glocial scatic acids and 4 drops of motherol-water (6:1 y/v). The minture was boiled for 8 minutes and kept overnight in a refrigerator. The mystalline product was filtered, weeked with cold otherol, efter recrystalline product was filtered, the map, of the derivative was found to be 210-15°, 14t. 25 , map. 215°.

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